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Choi et al.

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(54) **CRYSTAL STRUCTURE OF THE NANR AND MANNAC-6P COMPLEX, AND USES THEREOF**

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**C07K 14/28** (2006.01)

(52) **U.S. Cl.**

CPC ..... **G01N 33/56911** (2013.01); **C07K 14/28** (2013.01); **G01N 2400/00** (2013.01); **G01N 2500/04** (2013.01)

(58) **Field of Classification Search**

None

See application file for complete search history.

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NCBI Reference Sequence WP\_011081658.3, May 24, 2013.

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(57) **ABSTRACT**

The present invention relates to a three-dimensional structure of a complex explored by crystallization of the complex of NanR which is a key pathogenic regulatory protein of *Vibrio vulnificus* and ManNAc-6P which is a NanR regulator. Further, the present invention relates to a modified NanR protein, a polynucleotide encoding the protein, a vector including the polynucleotide, and a transformant including the vector. Furthermore, the present invention relates to a method for screening a substance regulating interaction between NanR and the transcriptional control region of nan operon which is a gene cluster regulated by NanR, or a substance regulating interaction between NanR and ManNAc-6P, by designing three-dimensional structure of the complex, and to an antibacterial composition including the screened substance.

**6 Claims, 26 Drawing Sheets**

FIG. 1

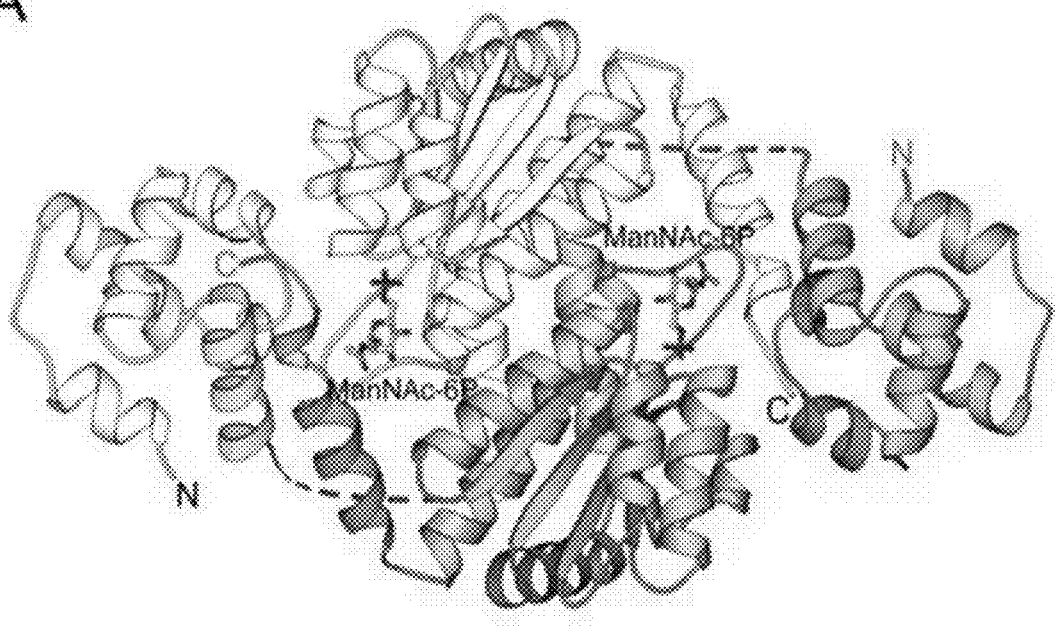
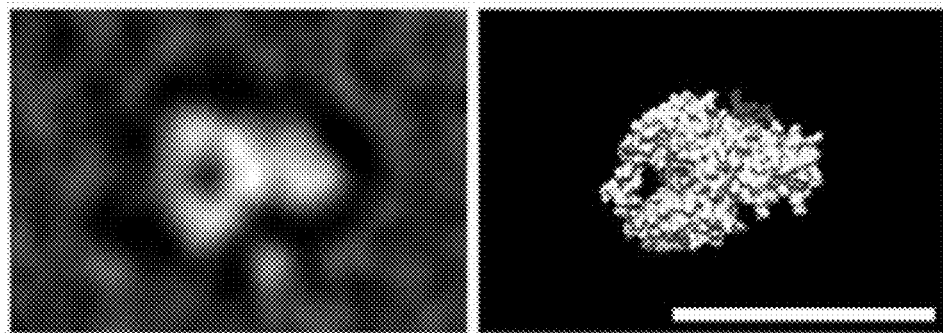
**A****B**

FIG. 2a

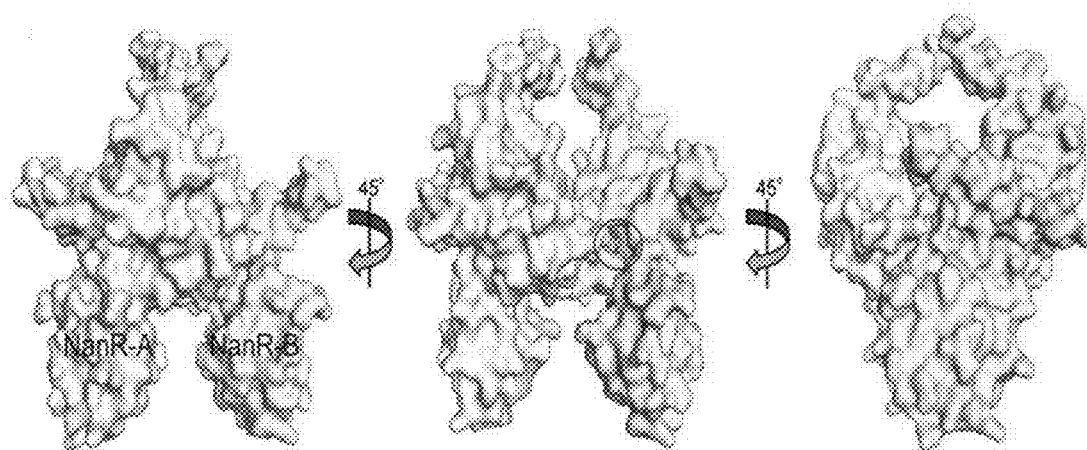


FIG. 2b

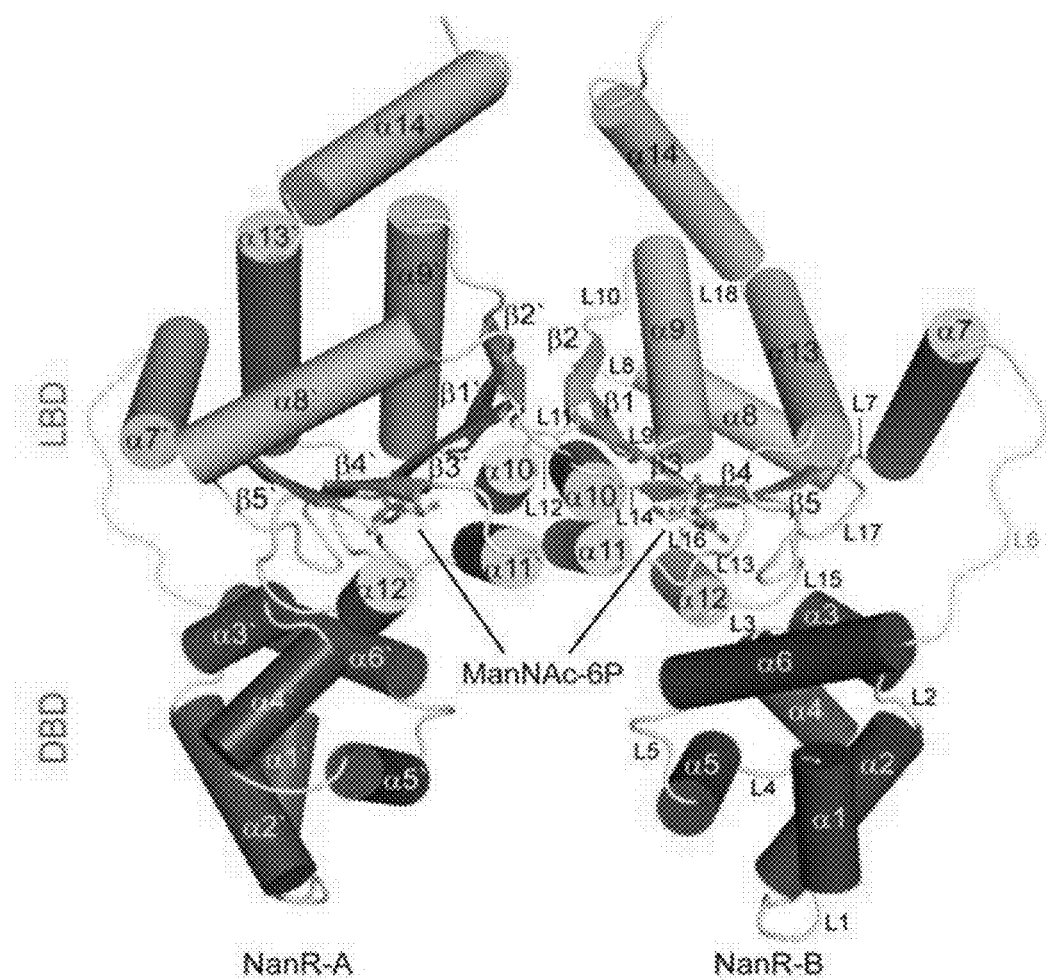


FIG. 3a

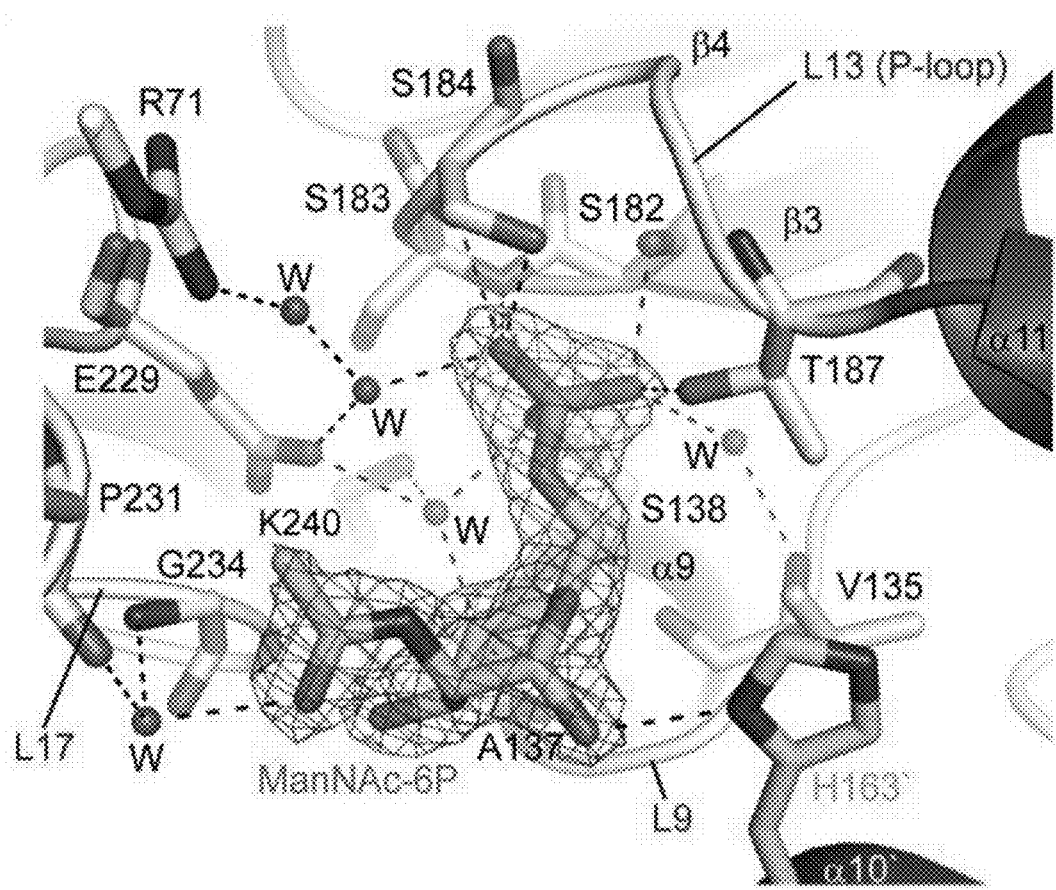


FIG. 3b

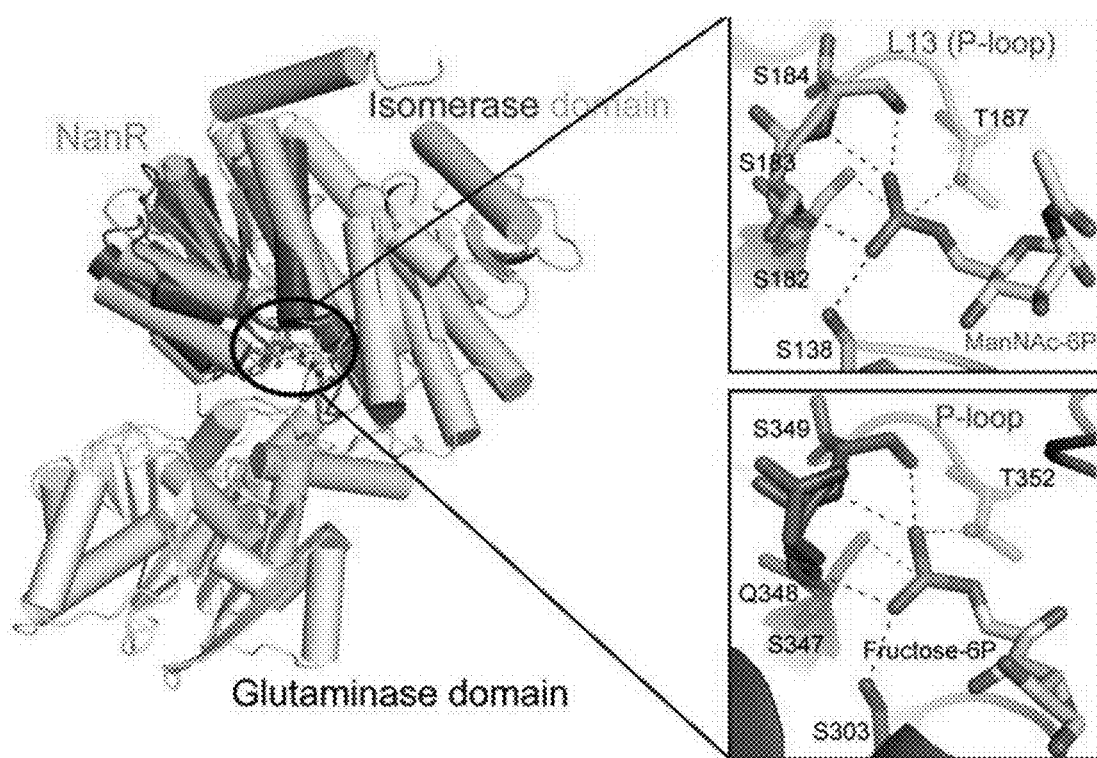


FIG. 3c

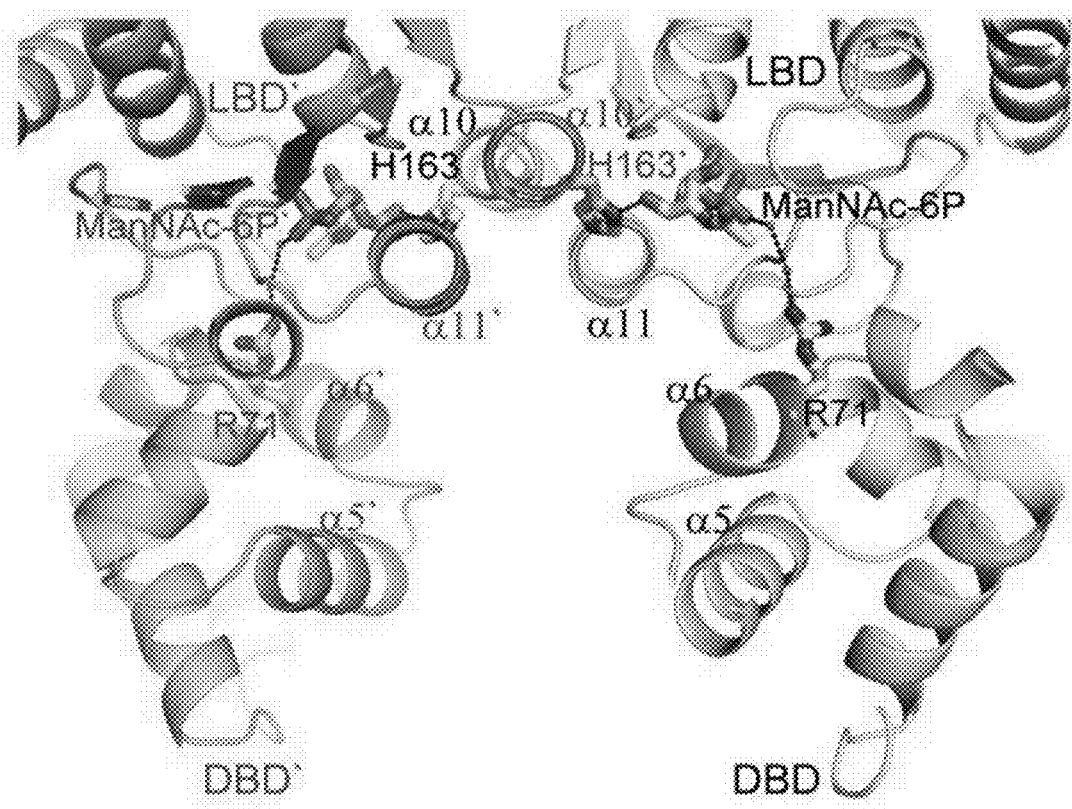


FIG. 3d

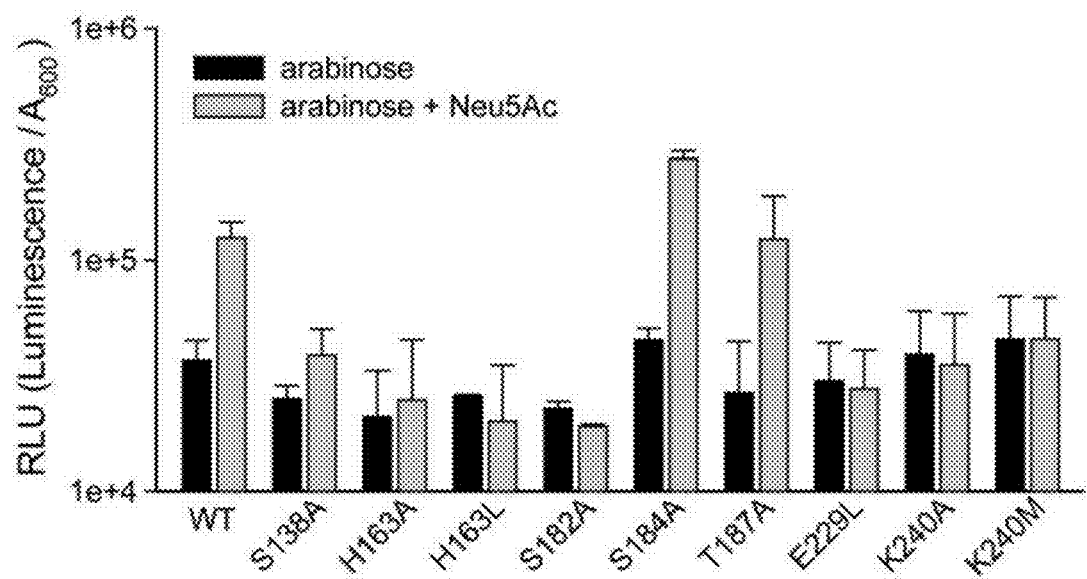


FIG. 4a

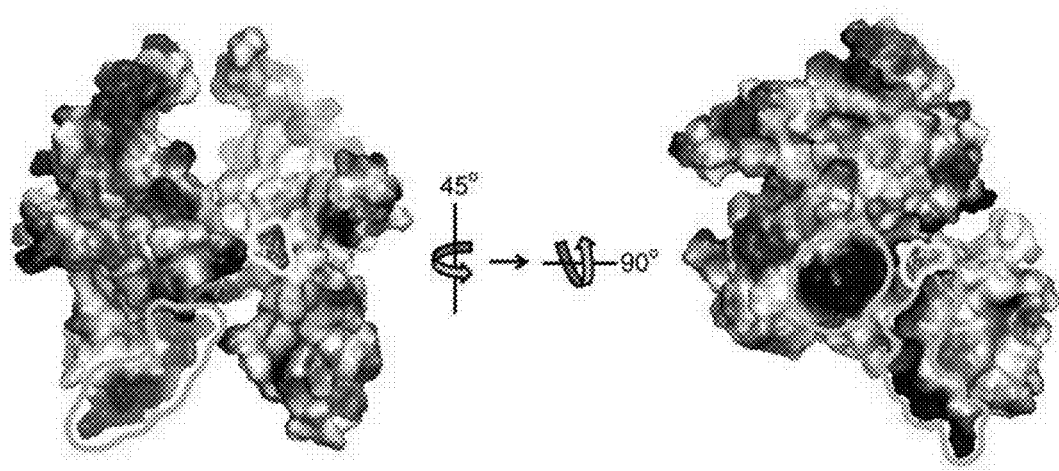


FIG. 4b

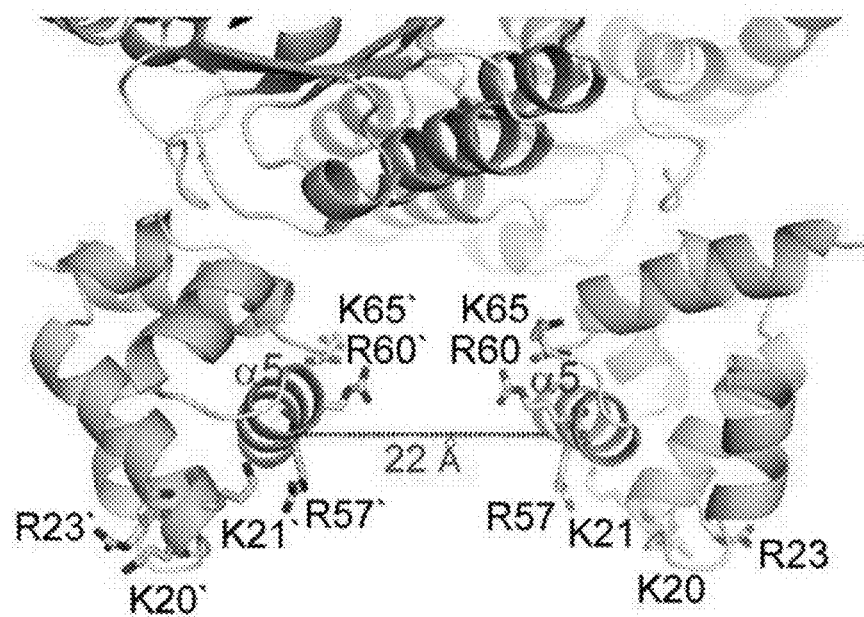


FIG. 4c

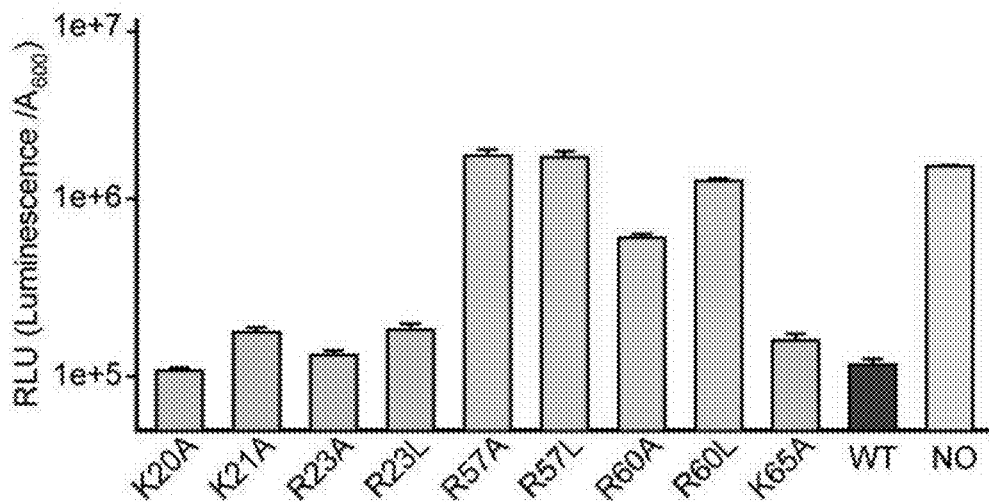


FIG. 4d

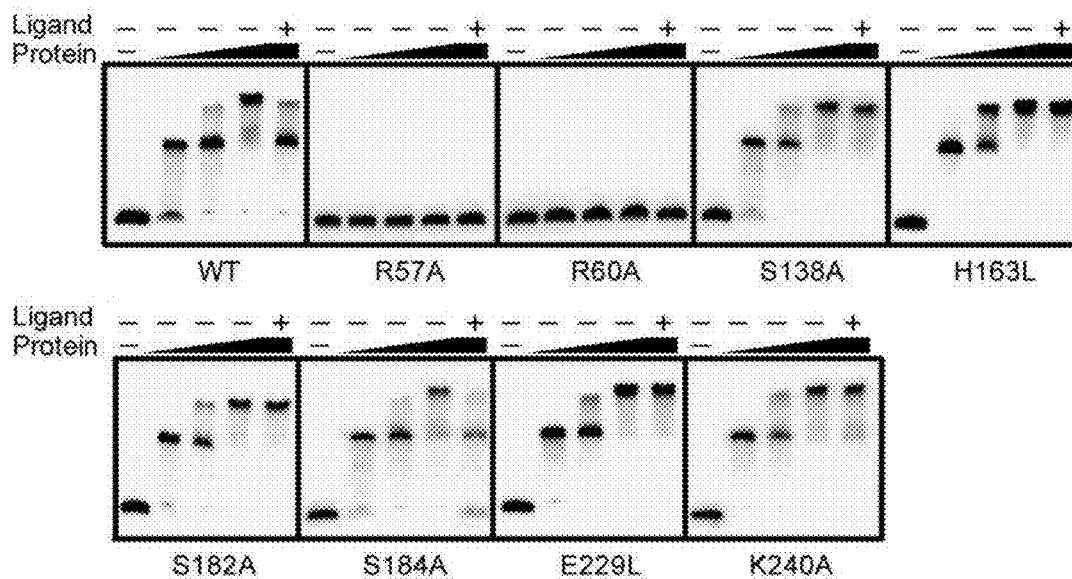


FIG. 4e

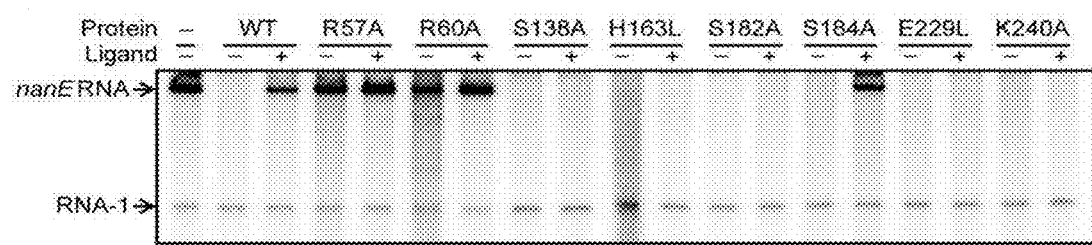


FIG. 4f

	n	K <sub>a</sub> (10 <sup>8</sup> M <sup>-1</sup> )	K <sub>d</sub> (μM)	ΔH (kcal mol <sup>-1</sup> )	TΔS (kcal mol <sup>-1</sup> )	ΔG (kcal mol <sup>-1</sup> )
NanR to DNA	1.34±0.001	71.60±3.36	1.40	54.90±0.33	62.87	-57.39
NanR/ligand to DNA	1.13±0.28	0.54±0.07	185.87	157.20±49.24	162.11	-4.91

FIG. 5

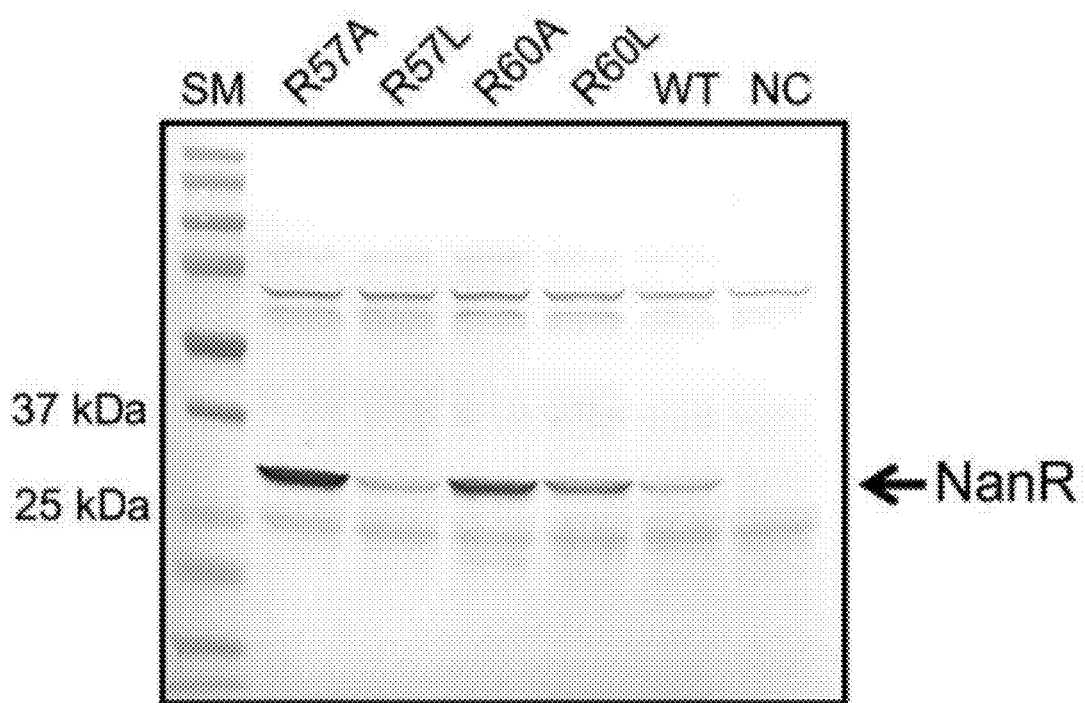


FIG. 6

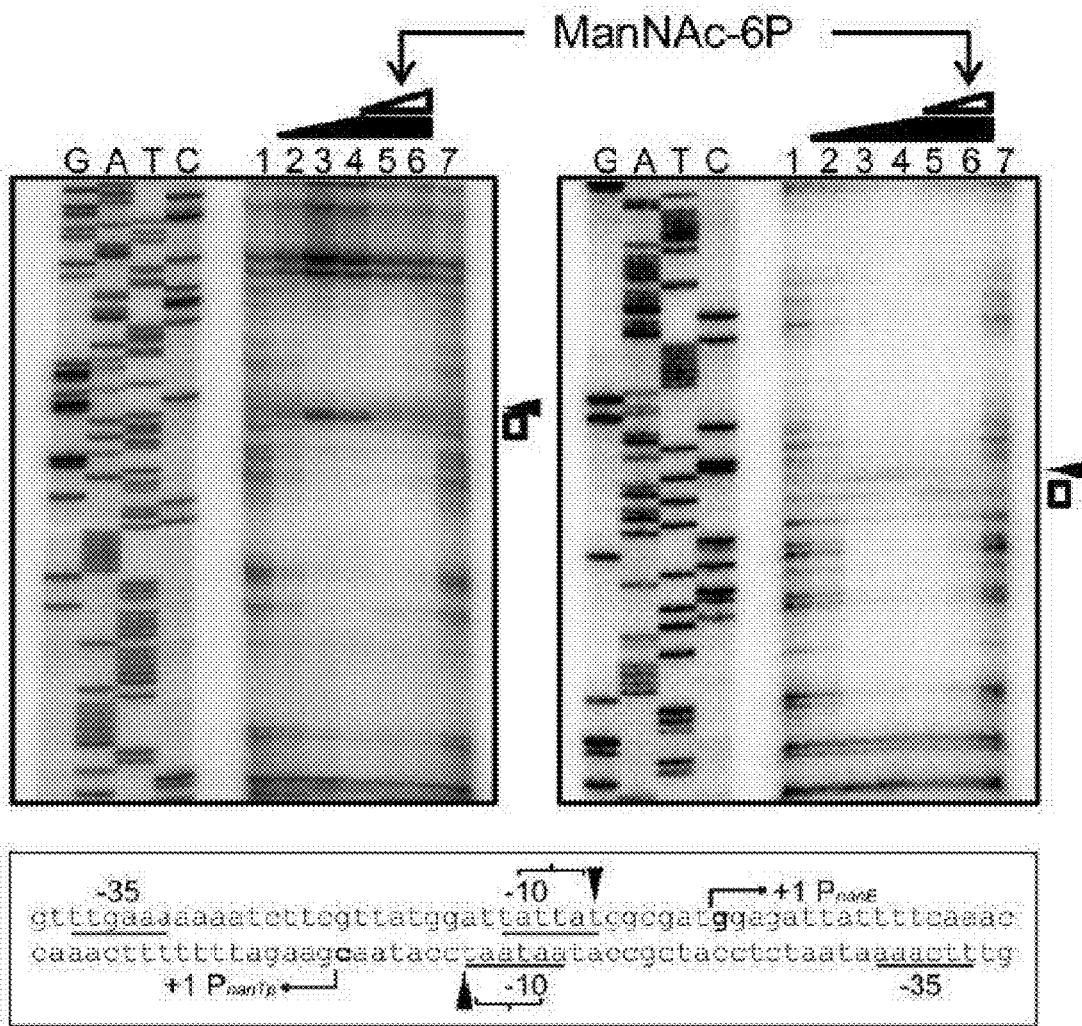


FIG. 7a

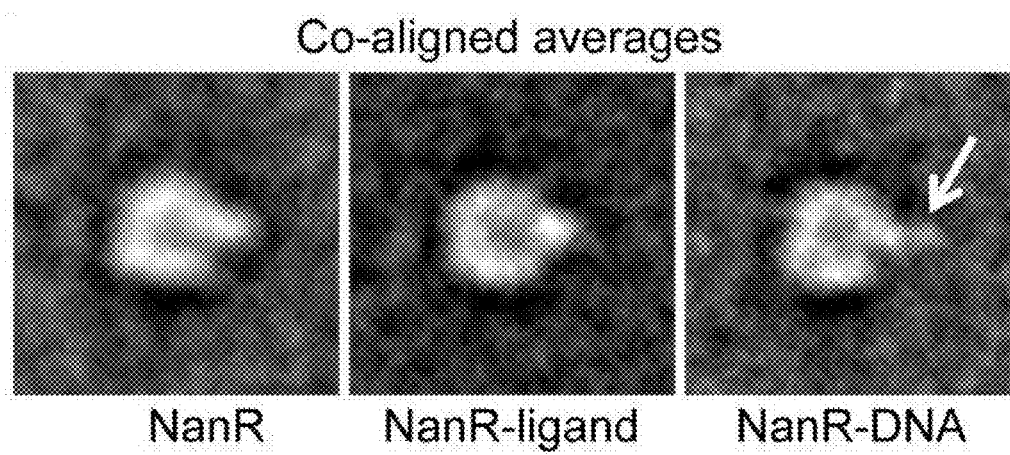


FIG. 7b

• Fitting of atomic models to averaged image

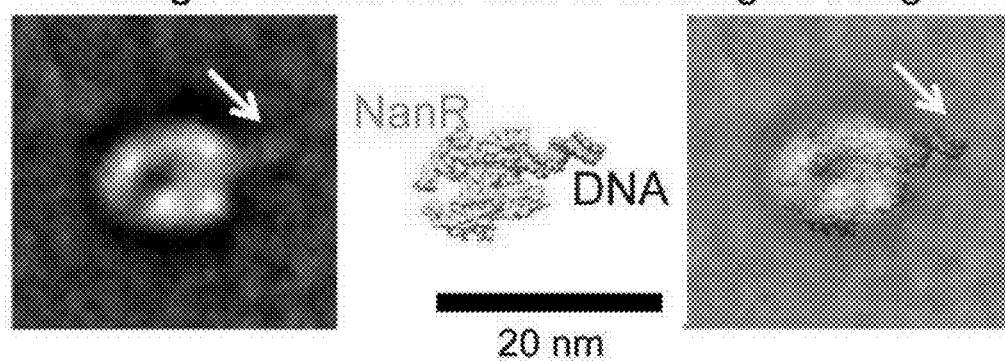


FIG. 7c

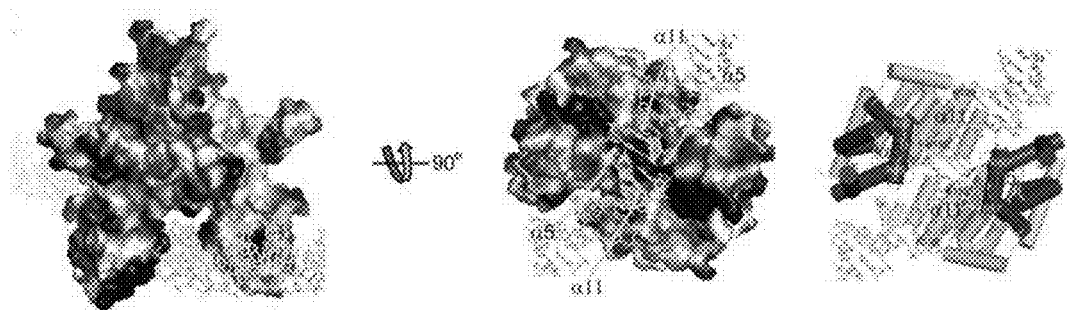


FIG. 8

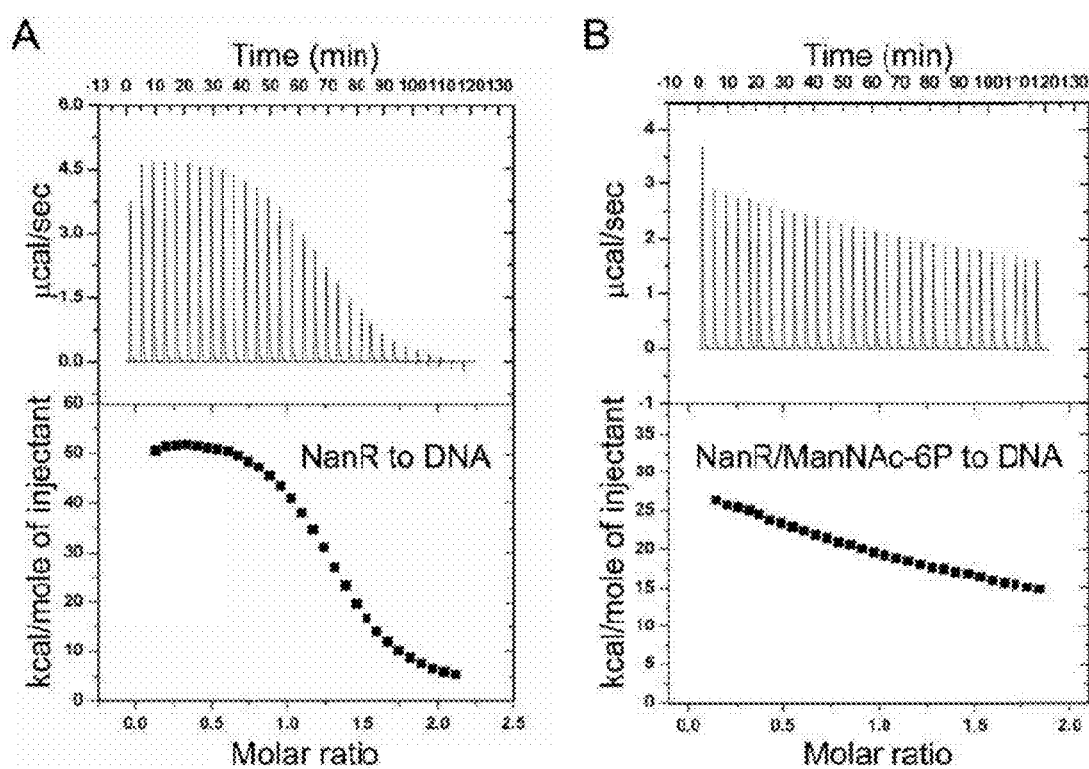


FIG. 9

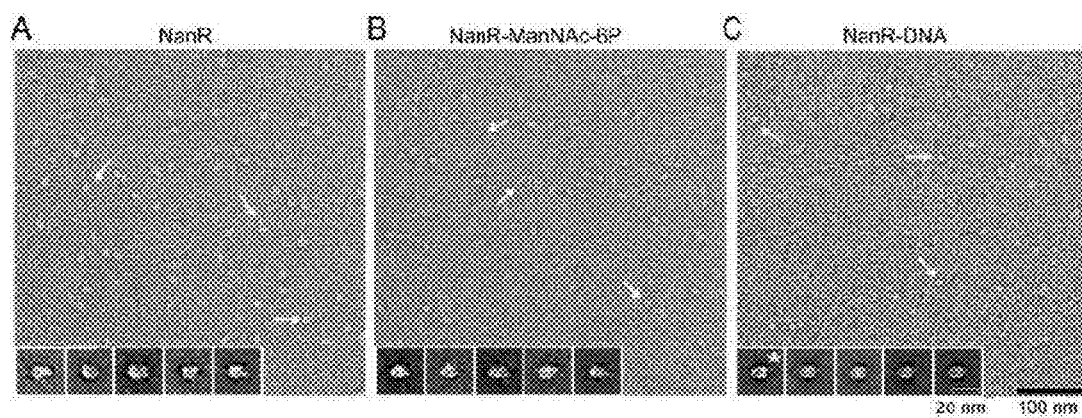


FIG. 10

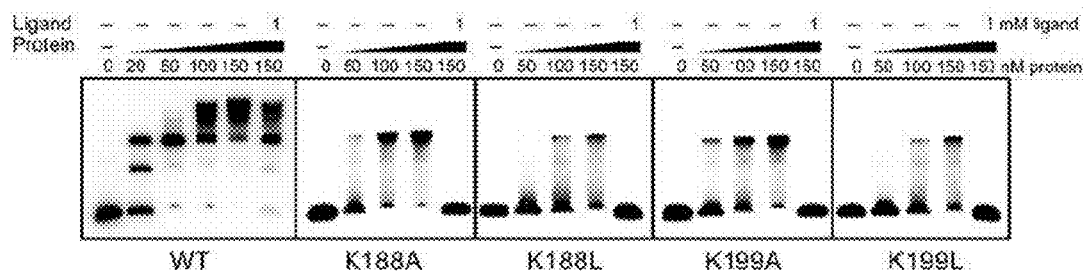


FIG. 11a

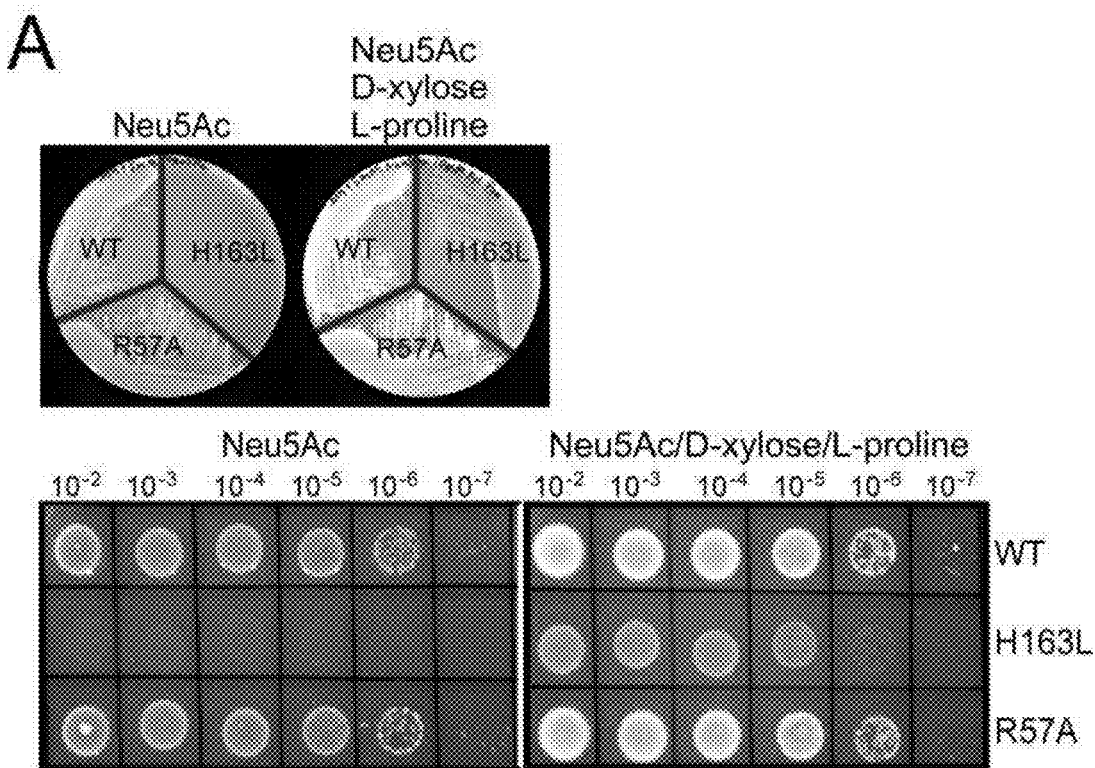


FIG. 11b

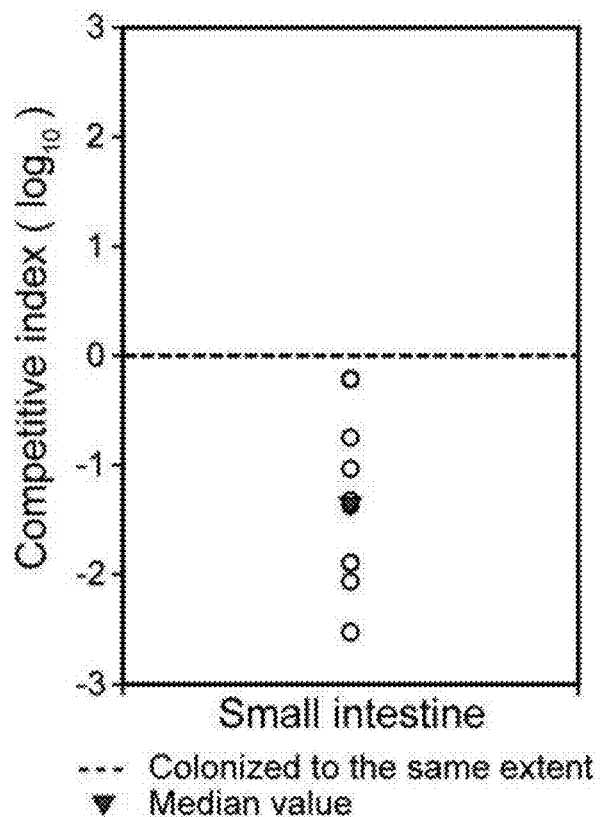


FIG. 11c

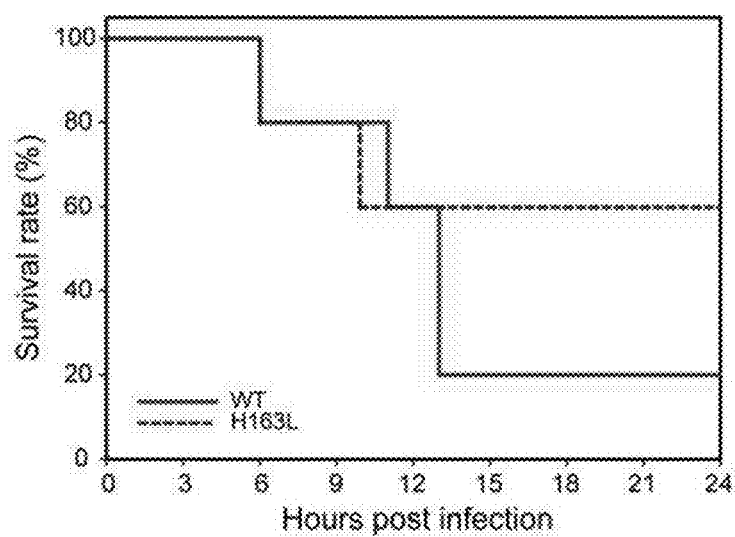


FIG. 11d

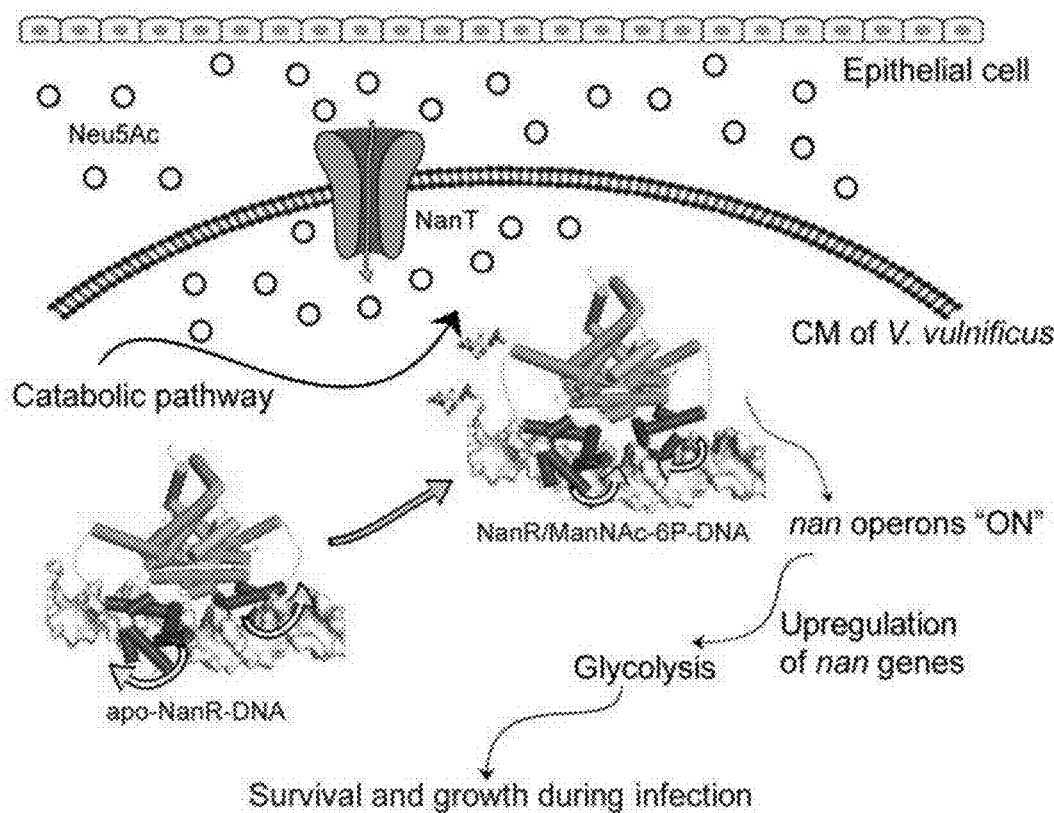


FIG. 12

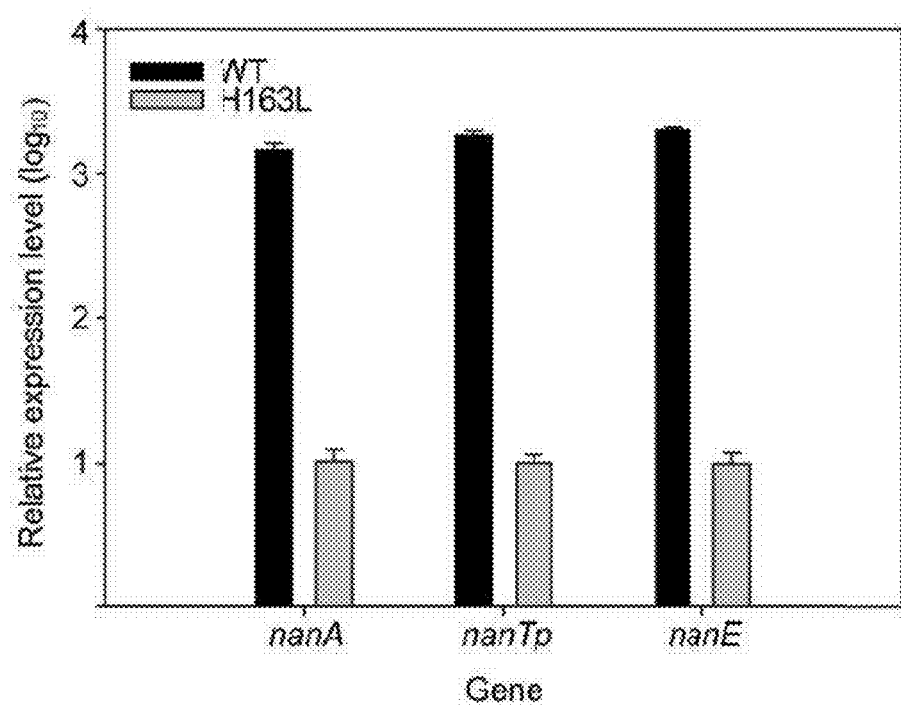
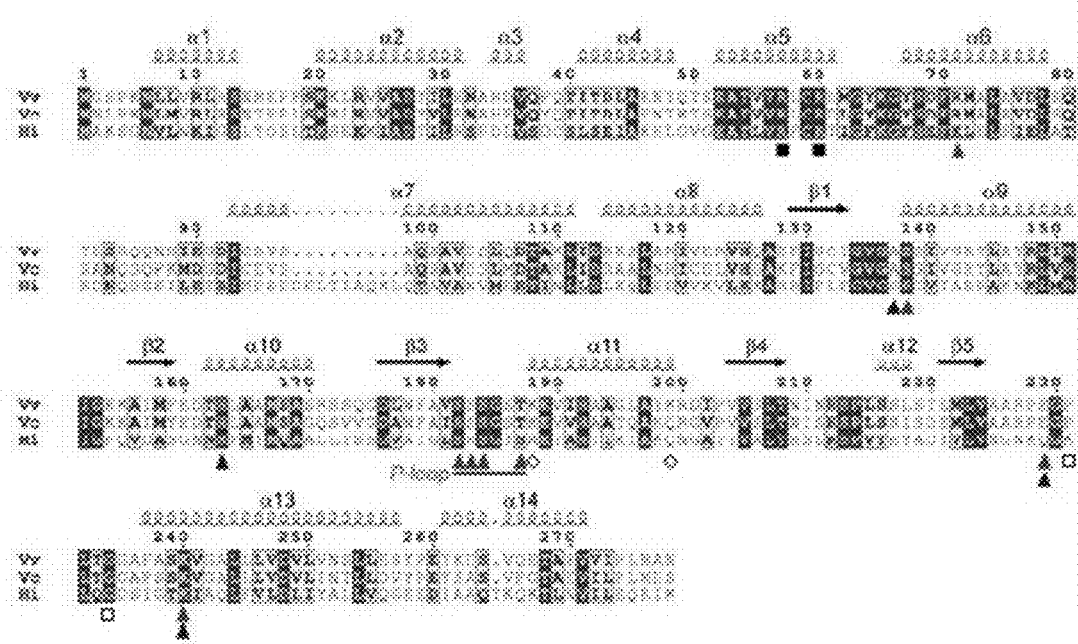


FIG. 13



## 1

**CRYSTAL STRUCTURE OF THE NANR AND  
MANNAC-6P COMPLEX, AND USES  
THEREOF**

**CROSS REFERENCE TO RELATED  
APPLICATIONS**

This application claims priority under 35 U.S.C. §119(a) to Korean Patent Application No. 10-2013-0071369, filed on Jun. 21, 2013, the disclosure of which is incorporated herein by reference.

**BACKGROUND OF THE INVENTION**

**1. Field of the Invention**

The present invention relates to a three-dimensional structure of crystallization of the complex of NanR which is a key pathogenic regulatory protein of *Vibrio vulnificus* and ManNAc-6P which is a NanR regulator. Further, the present invention relates to a modified NanR protein, a polynucleotide encoding the protein, a vector including the polynucleotide, and a transformant including the vector. Furthermore, the present invention relates to a method for screening a substance regulating interaction between NanR and the transcriptional control region of nan operon which is a gene cluster regulated by NanR, or a substance regulating interaction between NanR and ManNAc-6P, and an antibacterial composition including the screened substance.

**2. Description of the Related Art**

When the human body is infected with pathogenic bacteria, the pathogenic bacteria encounter competition with the gut intestinal flora for nutrients in the host (human body). Therefore, these pathogenic bacteria overcome the nutritionally adverse environment by utilizing alternative carbon sources in the gut, and endeavor to survive. The intestinal tract of the human body is protected by a mucus layer containing a glycosylated protein mucin that is composed of 85% carbohydrate. Pathogenic bacteria such as *Vibrio cholera* and *Vibrio vulnificus* (*V. vulnificus*) have most likely evolved elaborate systems for catabolic utilization of N-acetylneurameric acid (Neu5Ac), which is the most abundant sialic acids constituting mucin, as an alternative energy source.

When the human body is infected with these bacteria, these bacteria overexpress nan genes encoding enzymes essential for Neu5Ac catabolism and membrane transport proteins required for intracellular transport of Neu5Ac in the intestine so as to utilize Neu5Ac as an energy source, and thus they exert their pathogenicity through survival and growth, indicating that Neu5Ac catabolism of pathogenic microorganisms is directly correlated with their pathogenicity.

*V. vulnificus* is a pathogenic bacterium which usually enters the body through traumatic injury or ingestion of undercooked or contaminated sea food, and a life-threatening foodborne enteropathogen which causes septicemia in patients with liver disease or diabetes or in immunocompromised individuals, and the septicemia is associated with a mortality greater than 50% within 48 hours. Before entering the bloodstream, *V. vulnificus* survives and colonizes the small intestine. The present inventors recently demonstrated that NanR protein of *V. vulnificus* is a transcriptional repressor of the nan operon which is a cluster of genes encoding the transporter for intracellular absorption of Neu5Ac and essential catabolic enzymes for Neu5Ac, respectively. They also showed that N-acetylmannosamine 6-phosphate (ManNAc-6P), the catabolic intermediate of Neu5Ac, selectively

## 2

binds to NanR and induces expression of the nan genes. However, the molecular mechanisms underlying regulation of the nan genes by interaction between NanR protein and ManNAc-6P has not been clarified yet.

On the other hand, recent studies change their approach to development of therapeutic agents from a random approach to exploration of a number of therapeutic candidates to a new approach to development of therapeutic agents with improved target specificity and efficacy by exploration of key target proteins, investigation of three-dimensional structure and function of the target proteins, and designing and development of drug candidates through specific and efficient protein engineering. Therefore, to design and develop selective and specific drug candidates, it is essential that production and crystallization of a large amount of a highly pure, stable protein are conducted and then its three-dimensional structure is investigated. Three-dimensional structure of the NanR protein, in particular, a complex of the NanR protein and ManNAc-6P, for all its importance, has not been revealed yet, because it is difficult to crystallize in stable form.

Accordingly, the present inventors have made many efforts to investigate three-dimensional structure of the NanR protein. As a result, they prepared a crystal of NanR protein and its ligand ManNAc-6P complex, and investigated interaction between the NanR protein and its regulatory ligand at the atomic level with high resolution, thereby completing the present invention.

**SUMMARY OF THE INVENTION**

An object of the present invention is to provide a method for crystallizing a complex of NanR protein and N-acetylmannosamine 6-phosphate (ManNAc-6P).

Another object of the present invention is to provide a three-dimensional structure of the NanR protein and ManNAc-6P complex, and a crystal of SeMet-NanR protein and ManNAc-6P complex.

Still another object of the present invention is to provide a modified NanR protein, a polynucleotide encoding the protein, an expression vector including the polynucleotide, and a transformant introduced with the expression vector.

Still another object of the present invention is to provide a method for screening a substance regulating interaction between NanR and the transcriptional control region of nan operon which is a gene cluster regulated by NanR, by utilizing three-dimensional structure of the complex of NanR protein and ManNAc-6P.

Still another object of the present invention is to provide a method for screening a substance regulating interaction between NanR and ManNAc-6P, by utilizing three-dimensional structure of the complex of NanR protein and ManNAc-6P.

Still another object of the present invention is to provide an antibacterial composition including the screened substance.

**BRIEF DESCRIPTION OF THE DRAWINGS**

FIG. 1 shows symmetry mates in the dimer of NanR/ManNAc-6P complex (A) and an electron microscopic image of a functional dimeric form of NanR (B).

FIG. 2 shows an overall structure of the NanR/ManNAc-6P complex. FIG. 2a shows the binding structure of the dimer tilted by 45° and the ligand-binding site is indicated

by a circle; and FIG. 2b shows a DBD domain and a LBD domain, the  $\alpha$ -helices and  $\beta$ -sheets thereof, and the binding structure of ManNAc-6P.

FIGS. 3a to 3d show structural analysis of the interaction between NanR protein and ManNAc-6P. FIG. 3a is an electron density difference map showing the position of ManNAc-6P in the ligand-binding site, in which the residues critical for the interaction with NanR are shown together with ligands; FIG. 3b is the structure of NanR superimposed onto that of GlnS, which has an isomerase domain (upper) and a glutaminase domain (lower), in which the P-loop-binding ligand of each protein is shown; FIG. 3c shows that two NanR monomers form hydrogen bonds with each ligand; and FIG. 3d shows the results of an *E. coli* dual plasmid system assay, in which cells were cotransformed with a luciferase reporter gene fused to PnanTp and wild-type or mutant NanR, followed by incubation in the presence or absence of Neu5Ac, and the RLU (Relative luminescence unit) was calculated by dividing the luminescence by the A<sub>600</sub> of each strain and the data represent the mean±SD from at least three experiments.

FIGS. 4a to 4f show that the DNA-binding activity of NanR is regulated by ManNAc-6P. FIG. 4a is surface electrostatic potential showing the distribution of positively charged residues in the DNA binding domain (DBD), in which the positively charged residue is indicated by dark gray circle, and the ligand-binding site is indicated by light gray circle; FIG. 4b shows the position of the positively charged residues in the DBD domain of each NanR molecule; FIG. 4c shows the results of an *E. coli* dual plasmid system assay, in which cells were cotransformed with a luciferase reporter gene fused to PnanTp and wild-type NanR, mutant NanR, or empty vector (NO), followed by incubation, and the RLU (Relative luminescence unit) was calculated by dividing the luminescence by the A<sub>600</sub> of each strain, and the data represent the mean±SD from at least three experiments; FIG. 4d shows EMSA analysis of the interaction between the nanTp-nanE intergenic region and wild-type or mutant NanR protein in the absence or presence of ManNAc-6P ligand; FIG. 4e shows the results of in vitro transcription assay, in which the supercoiled pBS0921 plasmid containing PnanE was transcribed in the presence or absence of 100 nM wild-type or mutant NanR protein and 1 mM ManNAc-6P ligand, and the 370-bp PnanE-specific transcript and the vector-derived control transcript (RNA-1) are indicated; and FIG. 4f shows the effect of interaction between NanR and the transcriptional control region of nan operon, which was determined by using isothermal titration calorimetry.

FIG. 5 shows the result of immunoblot analysis for wild-type and mutant NanR expressions using an *E. coli* dual plasmid system, in which SM represents a size marker, and NC represents a negative control.

FIG. 6 shows the effect of ManNAc-6P ligand on binding of NanR to DNA (SEQ ID NO:15), in which DNaseI footprinting assay was performed by using NanR protein and 32P-labeled transcriptional control region of nan operon (299 bp) in the presence or absence of ManNAc-6P. After the addition of ManNAc-6P, the hypersensitive cleavage bands (black triangle, ▼) at the PnanE and PnanTp disappeared, and the neighboring regions were deprotected (white box and parentheses). Lanes 1 and 7 represent no addition of NanR; Lanes 2 and 3 represent 100 nM or 200 nM NanR treatment, respectively; Lanes 4 to 6 represent 400 nM NanR treatment; and Lanes 5 and 6 represent 0.1 mM or 1 mM ManNAc-6P treatment, respectively.

FIGS. 7a to 7c show the results of electron microscopy of NanR. FIG. 7a shows averaged images of apo-NanR, NanR/ManNAc-6P complex, apo-NanR/DNA complex; FIG. 7b shows the result of fitting the atomic models of NanR and DNA to the NanR-DNA complex; and FIG. 7c shows the structure of the NanR/DNA complex, which was modeled based on the electron-microscopic analysis and *in vivo* and *in vitro* investigations, in which R57 and R60 residues in  $\alpha$ 5 and K199 in  $\alpha$ 11 are found to be required for DNA binding.

FIG. 8 shows the results of isothermal titration calorimetry for interaction between DNA and NanR protein. (A) shows the result for interaction between DNA and apo-NanR protein, and (B) shows the result for interaction between DNA and NanR/ManNAc-6P complex, and the upper panel shows the raw data and the lower panel shows the result of analysis.

FIG. 9 shows electron microscopic images for negative staining of apo-NanR (A), NanR/ManNAc-6P complex (B) and apo-NanR/DNA complex (C), in which the white arrows indicate averaged images.

FIG. 10 shows the result of EMSA for examining the effect of K188 and K199 residues in all on binding of NanR to the transcriptional control region of nan operon, in which EMSA was carried out with increasing amount of each wild-type or mutant NanR protein, and 1 mM ManNAc-6P was further added to carry out the experiment.

FIGS. 11a to 11d show the effect of the ManNAc-6P-mediated regulation of NanR on bacterial pathogenesis. FIG. 11a shows that overnight cultures of wild-type, R57A, and H163L NanR mutant *V. vulnificus* strains were washed with PBS three times and serially diluted, the undiluted samples were streaked (left panel) onto a solid medium, and each serial dilution was spotted onto a medium supplemented with either Neu5Ac (lower left panel) as a sole carbon source or Neu5Ac, D-xylose, and L-proline (lower right panel), followed by incubation at 30° C. for 24 hours; FIG. 11b shows the results of a mouse intestine colonization competition assay using the wild-type and H163L NanR mutant *V. vulnificus* strain (n=10), in which each mouse was indicated by white circles and median values are indicated by triangles; FIG. 11c shows survival rates for 24 hours after mice were challenged with the wild-type and H163L NanR mutant *V. vulnificus* strain (4×10<sup>8</sup> CFU) (n=5 per group); and FIG. 11d shows molecular mechanism of Neu5Ac catabolism via the ManNAc-6P-mediated regulation of NanR, in which Neu5Ac is indicated by white circles and CM indicates cell wall.

FIG. 12 shows the result of analyzing the effect of H163L mutation of NanR protein on expression of nan genes in *V. vulnificus*, in which the wild-type or H163L NanR-expressing strain was cultured in the presence of Neu5Ac, total RNA was extracted, and the amount of the transcript was analyzed by qRT-PCR (normalization to 16S rRNA), and quantity of the transcript of H163L mutant was considered as 1 to calculate log<sub>10</sub>. The data represent the mean±SD from at least three experiments.

FIG. 13 shows conservation of *V. vulnificus*, *V. cholerae*, and *H. influenzae*-derived NanR sequences. Black squares (■) indicate residues essential for DNA-binding, gray triangles indicate residues involved in binding to the phosphate group of ManNAc-6P, empty squares (□) indicate residues involved in binding to the N-acetyl group of ligand, and diamonds (◇) indicate additional DNA-binding residues. Vv represents *V. vulnificus* UJ016 (gi:37676858) (SEQ ID NO:1); Vc represents *V. cholerae* O1 biovarE1 Tor str. N16961 (gi:15641775) (SEQ ID NO:16); and Hi represents *H. influenzae* Rd KW20 (gi:16272110) (SEQ ID NO:17).

T-COFFEE software was used for sequence comparison and ESPript software was used for visualization. These two softwares are available in ExPASy portal (<http://au>>--<<expasy.org/>).

#### DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

In one aspect to achieve the above objects, the present invention provides a method for crystallizing a complex of NanR protein and N-acetylmannosamine 6-phosphate (ManNAc-6P).

Specifically, the method for crystallizing the complex of NanR protein and ManNAc-6P includes the steps of (a) mixing the NanR protein having an amino acid sequence of SEQ ID NO. 1 with ManNAc-6P; and (b) crystallizing the mixture of step (a), but is not limited thereto.

As used herein, the term "NanR protein" is a transcriptional repressor protein suppressing expression of the genes in nan operon essential for catabolism of Neu5Ac (N-acetyl-neurameric acid) which is the most abundant sialic acid in the intestine, and it has an N-terminal DNA binding domain containing a helix-turn-helix(HTH) motif and a C-terminal domain binding to a ligand. In the present invention, the NanR protein includes all of the wild-type NanR protein and modified NanR protein. Further, the NanR protein of the present invention may be a modified NanR protein, in which methionine is substituted with selenomethionine (SeMet). Meanwhile, SeMet, as used herein, refers to an amino acid in which sulfur atom in methionine is replaced by selenium. SeMet shows behavior and chemical characteristics similar to those of methionine. SeMet is used to solve the phase problem for obtaining a three-dimensional crystal structure in X-ray crystallography, when no similar three-dimensional structure has been revealed so far. A crystallization method by substitution of methionine with SeMet for protein structural analysis is also called a SeMet method. Information about the amino acid sequence and nucleotide sequence of the NanR protein can be obtained from the known database such as NCBI GenBank. For example, the NanR protein may have the amino acid sequence of SEQ ID NO. 1. However, as long as a protein has the activity of the protein having the amino acid sequence of SEQ ID NO. 1, the protein having 80% or more homology, specifically 90% or more homology, more specifically 95% or more homology, and much more specifically 99% or more homology thereto may be also included without limitation. It is apparent that a protein having an amino acid sequence in which a part thereof is deleted, altered, substituted, or added is also included in the scope of the present invention.

In one embodiment of the present invention, to solve the phase problem of the crystallized protein, SeMet-NanR was prepared by substitution of SeMet for methionine constituting the protein, and mixed with ManNAc-6P, followed by crystallization. In order to obtain the SeMet-substituted NanR protein, the methionine auxotroph *E. coli* B834 (DE3) strain (Novagen) was grown in a minimal medium supplemented with 50 mg/ml SeMet. The SeMet-substituted NanR protein was purified and obtained from the culture by addition of 5 mM methionine to all buffers.

Moreover, the NanR protein may be derived from, but is not limited to, *V. vulnificus*, *Escherichia coli*, *Haemophilus influenzae* or *V. cholerae*.

Meanwhile, the present inventors recently revealed that *V. vulnificus* NanR protein is a transcriptional repressor of nan operon which is a cluster of genes encoding transporters for intracellular absorption of Neu5Ac and enzymes essential

for Neu5Ac catabolism, respectively. In addition, they also demonstrated that the catabolic intermediate of Neu5Ac, ManNAc-6P selectively binds to NanR to induce expression of nan genes (BS Kim et al., THE JOURNAL OF BIOLOGICAL CHEMISTRY, Vol. 286, No. 47, pp. 40889-40899, 2011 Nov. 25). With respect to the objects of the present invention, the NanR protein may be a protein binding to the transcriptional control region of nan operon, and specifically, the NanR protein may be a protein that binds to the nan operon promoter to suppress expression of nan operon, but is not limited thereto. The NanR protein may be also a protein binding to ManNAc-6P, and its binding to ManNAc-6P may cause conformational change, leading to a reduction in binding affinity with the transcriptional control region of nan operon, but is not limited thereto.

As used herein, the term "modified NanR protein" means a protein which is prepared by substituting, inserting, deleting or modifying one or more amino acids in the wild-type NanR protein, and with respect to the objects of the present invention, the modified NanR protein may be a protein which is modified to have increased or reduced binding affinity for the transcriptional control region of nan operon or for the ManNAc-6P ligand, but is not limited thereto.

The modified NanR protein is preferably a protein having modifications of one or more selected from the group consisting of lysine at position 20 (Lys20), lysine at position 21 (Lys21), arginine at position 23 (Arg23), arginine at position 57 (Arg57), arginine at position 60 (Arg60), lysine at position 65 (Lys65), serine at position 138 (Ser138), histidine at position 163 (His163), serine at position 182 (Ser182), serine at position 184 (Ser184), threonine at position 187 (Thr187), glutamic acid at position 229 (Glu229), and lysine at position 240 (Lys240) in the amino acid sequence of SEQ ID NO. 1 of NanR protein, for example, a protein having substituted amino acids, and more preferably, a protein in which Lys20 is substituted with alanine, Lys21 is substituted with alanine, Arg23 is substituted with alanine or leucine, Arg57 is substituted with alanine or leucine, Arg60 is substituted with alanine or leucine, Lys65 is substituted with alanine, Ser138 is substituted with alanine, His163 is substituted with alanine or leucine, Ser182 is substituted with alanine, Ser184 is substituted with alanine, Thr187 is substituted with alanine, Glu229 is substituted with leucine, or Lys240 is substituted with alanine or methionine, but is not limited thereto. In addition to the above modified proteins, it is apparent that a modified NanR protein having 70% or more homology, preferably 80% or more homology, more preferably 95% or more homology, and most preferably 98% or more homology to the amino acid sequence of SEQ ID NO. 1 is also included in the scope of the present invention, as long as its binding ability to the transcriptional control region of nan operon or to the ManNAc-6P is controlled, compared to the wild-type NanR protein. The modified protein of the present invention may be a protein that can undergo crystallization required for 3D structural analysis of the protein.

In one embodiment of the present invention, a modified NanR protein was prepared by substituting Lys20 with alanine, Lys21 with alanine, Arg23 with alanine or leucine, Arg57 with alanine or leucine, Arg60 with alanine or leucine, Lys65 with alanine, Ser138 with alanine, His163 with alanine or leucine, Ser182 with alanine, Ser184 with alanine, Thr187 with alanine, Glu229 with leucine, or Lys240 with alanine or methionine in the NanR protein having the amino acid sequence of SEQ ID NO. 1, and its binding affinity for the transcriptional control region of nan operon or the ManNAc-6P was examined.

As used herein, the term “ManNAc-6P” is, also called acetylmannosamine-6 phosphate, a catabolic intermediate of N-acetylneurameric acid which constitutes glycoproteins, and synthesized by action of N-acylmannosamine kinase. That is, it is a catabolic intermediate of N-acetylneurameric acid which is the most common sialic acid found in nature, such as animal cell membranes, glycoproteins and glycolipids, or bacterial cell walls, and ManNAc-6P is present from virus to animals. ManNAc-6P has a chemical formula of C<sub>8</sub>H<sub>16</sub>NO<sub>9</sub>P. In the present invention, ManNAc-6P binds with the NanR protein, which causes conformational change, leading to a reduction in binding affinity of NanR protein with the transcriptional control region of nan operon, but is not limited thereto. In particular, the ManNAc-6P may binds to the C-terminal ligand of the NanR protein. Eventually, ManNAc-6P functions to reduce the nan gene transcriptional repressor activity of the NanR protein, thereby inducing expression of nan operon genes.

As used herein, the term “complex of NanR protein and ManNAc-6P” means a complex formed by interaction between NanR protein and ManNAc-6P. Preferably, it means a complex formed by interaction between arginine residue at position 71 (Arg71) of a DNA binding domain (DBD) and alanine residue at position 137 (Ala137), Ser138, His163, Ser182, serine residue at position 183 (Ser183), Ser184, Thr187, Glu229, proline residue at position 231 (Pro231), glycine at position 234 (Gly234), and Lys240 of a ligand binding domain (LBD) of NanR protein and ManNAc-6P via a hydrogen bond or a water-mediated hydrogen bond, but is not limited thereto.

In one specific embodiment of the present invention, the binding structure between NanR protein and ManNAc-6P was examined by analyzing the structure of the crystal prepared in the present invention. In detail, ManNAc-6P binds to L9, L13, and L17 loops of NanR LBD domain (FIG. 3a). L13 crosses over and embraces the phosphate group of ManNAc-6P (FIGS. 3a and 3b); the phosphate oxygen atoms form hydrogen bonds with the side chains of S182, S184, and T187, and with the backbone amide of S183 in the P-loop (FIG. 2b, upper panel). The side chain hydroxyl group of S138 in loop L9 is also hydrogen-bonded to a ManNAc-6P phosphate oxygen atom (FIGS. 3a and 3b). The hydroxyl group at position O4 of the sugar ring forms a hydrogen bond with the A137 backbone amide. A hand-in-hand interaction between the two NanR monomers is formed by hydrogen bonds between the hydroxyl group at position O1 of the sugar ring in each NanR monomer and the nitrogen atom in the imidazole ring of each H163 (FIGS. 3a and 3b). This structure is critical for the conformational change of the NanR dimer and delivery of the signal to the nan operon genes when ligand-binding occurs. Furthermore, the phosphoryl group of the ligand forms a water-mediated hydrogen bond with the side chain amino group of R71 on  $\alpha$ 6 in the DBD domain (FIGS. 3a and 3c). P231 and G234 form water-mediated hydrogen bonds with the carbonyl oxygen atom of the N-acetyl group (FIG. 3a). In addition, E229 and K240 form water-mediated hydrogen bonds with the sugar and phosphate oxygen atoms of ManNAc-6P (FIG. 3a). These interactions may enable ligand-mediated relocation of the NanR dimer and influence its interaction with the transcriptional control region of nan operon.

In the crystallization method, the mixing step may be carried out by mixing NanR protein and ManNAc-6P in a molar ratio of 1:10 to 1:200, but is not limited thereto. In one embodiment of the present invention, NanR protein and ManNAc-6P were mixed in a molar ratio of 1:100, resulting in successful crystallization.

The NanR protein may be a protein that is separately expressed or obtained by purification of naturally occurring protein. The purification may be a purification method well known in the art. Therefore, the crystallization method may include the step of purifying the NanR protein before the mixing step of (a), and purification of the protein may be carried out by a known purification method such as affinity chromatography.

For example, the protein may be isolated from the medium, in which host cells are grown, by the conventional chromatographic method including immunoaffinity chromatography, receptor affinity chromatography, hydrophobic interaction chromatography, lectin affinity chromatography, size exclusion chromatography, cation or anion exchange chromatography, high performance liquid chromatography (HPLC), reverse phase HPLC or the like. In other purification method, the NanR protein is fused with a specific tag, label, or chelating moiety, and the fusion protein is recognized by a specific binding partner or agent. The purified protein can be cleaved to yield the desired protein, or can be left as an intact fusion protein. A form of the desired polypeptide having additional amino acid residues as a result of the cleavage process of the fusion protein may be produced.

As used herein, “undergoing crystallization” or “having crystallinity” means that in order to prepare a protein in the form suitable for X-ray analysis of three-dimensional structure, a modification is introduced into the protein molecule so as to form solid particles having a uniform shape and size from a uniform liquid or to further stabilize the crystal of the protein. Three-dimensional structure of a protein is very important for the understanding of *in vivo* actions of the protein and development of therapeutic drugs. That is, understanding of arrangement and three-dimensional structure of atoms constituting a protein as a macromolecule makes it possible to analyze three-dimensional structure of the complex of NanR protein and ManNAc-6P and to provide a platform for development of new drugs for inhibiting interaction between NanR protein and ManNAc-6P. However, it was very difficult to demonstrate three-dimensional structure of a protein or a complex of a protein and a ligand, because a crystal structure of the corresponding complex should be first prepared in order to analyze three-dimensional structure of the complex of NanR protein and ManNAc-6P. In addition, acquisition of a stable crystal depends on the protein and the ligand, and in particular, crystallization conditions vary depending on the protein and the ligand.

In the present invention, the present inventors mixed the NanR protein with ManNAc-6P in a molar ratio of 1:100 in order to obtain the crystal complex of NanR protein and ManNAc-6P.

The crystallization step of (b) may be carried out in a solution containing 1 to 20% PEG 2000 MME, 0.01 to 1 M ammonium sulfate, 0.01 to 2 M sodium formate, 0.5 to 6% low molecular polyglutamic acid (PGA-LM), and 0.01 to 1 M sodium acetate, and in particular, in a solution containing 10 to 20% PEG 2000 MME, 0.05 to 0.2 M ammonium sulfate, 0.1 to 0.5 M sodium formate, 1 to 5% low molecular polyglutamic acid, and 0.1M sodium acetate at pH 5.0 to pH 5.5. In the crystallization step, the mixture was incubated on ice for 1 to 24 hours.

In one embodiment of the present invention, crystallization trials of the purified NanR protein performed using the sitting drop vapor-diffusion method at 21° C. were unsuc-

cessful. However, the crystals were obtained when NanR and ManNAc-6P were mixed in a molar ratio of 1:100 and incubated on ice for 2 hours.

The crystallization of the complex was optimized under the following conditions: 10% PEG 2,000 MME, 0.1 M ammonium sulfate, 0.3 M sodium formate, 3% PGA-LM, and 0.1 M sodium acetate (pH 5.0 to 5.5). Crystals appeared within a day and were grown for a further 5 days for diffraction experiments. The crystals were transferred to a cryoprotectant solution containing 10% PEG 2,000 MME, 0.1 M ammonium sulfate, 0.3 M sodium formate, 3% PGA-LM, 0.1 M sodium acetate (pH 5.5), and 30% glycerol, and then placed immediately in a -173° C. nitrogen gas stream.

The crystallization of step (b) may be carried out by a variety of known crystallization methods, specifically, by a vapor diffusion method. The vapor diffusion method may be a sitting drop vapor diffusion method or a hanging-drop vapor diffusion method, and more specifically, sitting drop vapor diffusion method, but is not limited thereto.

As used herein, the term "sitting drop vapor diffusion method" refers a crystallization method, in which when a microdrop of mother liquor and a much larger reservoir solution exist separately in a closed system, transport of either water or other volatile agent occurs between them, leading to a supersaturated state of protein, and in such a thermodynamically metastable state, proteins are precipitated depending on the precipitant. While the protein precipitation slowly occurs, stable crystals are formed and the precipitant functions to lower the solubility of the concentrated protein solution, and proteins congregate to form crystals in order to reduce an adsorption layer around protein molecules. The reservoir solution contains a mixture of the precipitant, buffer, and detergent at different concentrations. Droplets are usually formed by mixing the protein solution with the reservoir solution of various conditions at a ratio of 1:1, and the droplets thus formed are placed on a microbridge, and sealed. At this time, there is a difference in the concentration between the proteins in the droplets and the reservoir solution, and thus the proteins do not exist as crystals at the beginning. They are equilibrated while sealed, and crystals are formed under the specific conditions by the above described principle. In the sitting drop vapor diffusion method, the type and proper concentration of the salt, the buffer and the surfactant as well as the precipitant in the reservoir solution, pH of the solution, and the experimental temperature vary depending on the type of protein, and in some cases, they become very important factors in crystal formation of proteins.

As used herein, the term "hanging-drop vapor diffusion method" is one of protein crystallization methods, which provides crystals having a size sufficient for protein structural analysis. In the hanging-drop vapor diffusion method, a reagent containing a sample and a pure liquid reagent are placed on the top of the reservoir under vapor equilibration. To achieve equilibrium of the sample having a lower reagent concentration than the reservoir, water contained in the sample eventually ends up in the reservoir. Water contained in the sample is removed until the concentration is approximately the same as that in the liquid reagent, and finally, protein crystals reaching the equilibration can be obtained.

In one embodiment of the present invention, the sitting drop vapor diffusion method was used to obtain complex crystals of NanR protein and ManNAc-6P (Example 8).

In another aspect, the present invention provides a crystal of NanR protein and ManNAc-6P complex or a crystal of SeMet-NanR protein and ManNAc-6P complex.

The NanR protein, ManNAc-6P, complex thereof, and the SeMet-NanR protein are the same as described above.

There are many different methods for analyzing a protein's crystal structure, and two main methods are NMR spectroscopy and X-ray crystallography. NMR Spectroscopy is based on the principle of predicting a distance between particular atoms in a molecule by analyzing signal changes due to chemical factor which can be detected in the NMR spectrum of a molecule. Data of chemical shift obtained by the NMR test is analyzed to obtain a set of the distances between labeled atoms in one protein, and a model or a set of models satisfying information about all distances determined by the experiment is produced. Thus, there is a disadvantage of requiring collection and analysis of a large amount of data. Meanwhile, X-ray crystallography is based on the principle of obtaining the result by analyzing x-ray scattered by electron cloud surrounding an atom of the crystal in an x-ray generator. X-ray diffraction patterns from protein crystals are regular because the individual protein molecules are arranged in a regular lattice. Based on this principle, X-ray crystallography is a method of analyzing a protein structure by producing an electron density of the protein using x-ray scattered and reflected from the protein crystals. However, there is a disadvantage of requiring pure protein samples and protein crystallization. In the present invention, crystallization trials of the NanR protein alone were unsuccessful, but the crystals were obtained when a complex of the NanR protein and its ligand ManNAc-6P was prepared.

In the present invention, to provide a three-dimensional crystal structure of NanR protein and ManNAc-6P, diffraction patterns were obtained using an x-ray image plate, and phase information was obtained by multiple anomalous dispersion (MAD) using Se-Met (selenium-methionine). An electron density map was obtained from the x-ray diffraction patterns and phase information, and atomic coordinates were derived therefrom so as to obtain the three-dimensional structure.

Preferably, the crystal of NanR protein and ManNAc-6P complex of the present invention has a space group of  $P3_{1}21$  and a unit-cell dimension of  $a=109.21\pm5$  Å,  $b=109.21\pm5$  Å and  $c=82.47\pm5$  Å,  $\alpha=\beta=90^\circ$  and  $\gamma=120^\circ$ , and specifically,  $a=109.21$  Å,  $b=109.21$  Å and  $c=82.47$  Å,  $\alpha=\beta=90^\circ$  and  $\gamma=120^\circ$ , and has an amino acid sequence of SEQ ID NO. 1, but is not limited thereto (Table 2).

Further, the complex crystal of SeMet-NanR protein and ManNAc-6P of the present invention has a space group of  $P3_{1}21$  and a unit-cell dimension of  $a=109.84\pm5$  Å,  $b=109.84\pm5$  Å and  $c=83.38\pm5$  Å,  $\alpha=\beta=90^\circ$  and  $\gamma=120^\circ$ , and specifically,  $a=109.84$  Å,  $b=109.84$  Å and  $c=83.38$  Å,  $\alpha=\beta=90^\circ$  and  $\gamma=120^\circ$ , and the NanR protein has the amino acid sequence of SEQ ID NO. 1, but is not limited thereto (Table 2).

As used herein, the term "space group" means the symmetry of a unit cell of a crystal, and combinations of symmetry elements form symmetry groups. This space is used interchangeably with the space group.

As used herein, the term "unit-cell dimension" is also called lattice parameter, and the unit-cell is the simplest minimum repeating unit constituting the space group, and defined by three crystallographic axes, the lengths of the three vectors ( $a$ ,  $b$ ,  $c$ ) and the inter-axial angles ( $\alpha$ ,  $\beta$ ,  $\gamma$ ).

The phase information can be obtained by multiple isomorphous replacement, multiwavelength anomalous dispersion, molecular replacement or the like. First, multiple isomorphous replacement is a technique of obtaining the phase information by replacing crystals with heavy metals,

and collecting and analyzing the data. Second, multiwavelength anomalous dispersion is a technique of obtaining phase information by collecting data using the anomalous dispersion at different wavelengths, in which a specific metal or atom in the crystal is used instead of heavy metals. That is, without the need for data collection of many crystals, data can be easily obtained from one crystal using selenium atom by replacing the amino acid methionine with selenomethionine (Se-Met) using a molecular biological method. However, this method has a disadvantage that data should be obtained from radiation beam. Third, molecular replacement is a method of solving the phase problem from the known similar structure, and it is widely used as the number of known structures is increasing. Data is collected from each structure, and then refinement is conducted to fit our model against the data. This procedure is performed using the known programs (CCP4, Coot, Quanta, CNS, etc.), and standardization of each angle, bond length, etc. is required. In this process, a procedure for fitting the model to the obtained electron density map by computer performance and by eye is repeatedly performed. In the analysis step after refinement of the structure, a lot of information can be derived from the structure with interpretation. In this analysis step, the mechanism of action can be studied, based on the structure. The studies on the correct mechanism of action provide information needed for development of new drugs. Further, the directly related residues can be identified through the structure of the complex of the protein and its regulator, and therefore, important information is provided for the next step for studying the regulator.

In one embodiment of the present invention, diffraction data for the crystal of the present invention were collected at 1.9 Å resolution. Single wavelength anomalous diffraction data for the SeMet-substituted crystals were collected at 2.4 Å resolution. All data were processed with HKL2000 software package. The structure of the NanR/ManNAc-6P complex was determined by analyzing the anomalous signals from Se atoms with the SOLVE program. Density modification and subsequent automated model building were performed using the RESOLVE program. The complex crystal structure was solved at 1.9 Å resolution by molecular replacement with the MOLREP program using the partially refined model of the SeMet crystal. The complex crystal structure was revised using the COOT program and refined using the REFMAC5. The atomic coordinates and structure factor amplitudes of NanR protein/ManNAc-6P obtained by the method have been deposited in the PDB (Protein Data Bank) with accession code of 4IVN. Further, information about this crystal is shown in Table 2.

In still another aspect, the present invention provides a modified NanR protein, a polynucleotide encoding the protein, an expression vector including the polynucleotide, and a transformant introduced with the expression vector.

The modified NanR protein is the same as described above.

The expression vector including the polynucleotide encoding the modified NanR protein provided in the present invention may be, but is not particularly limited to, a vector capable of replicating and/or expressing the polynucleotide in eukaryotic or prokaryotic cells including mammalian cells (e.g., human, monkey, rabbit, rat, hamster, mouse cells, etc.), plant cells, yeast cells, insect cells, or bacteria cells (e.g., *E. coli*, etc.), preferably, a vector which is operably linked to a suitable promoter to express the polynucleotide in the host cells and includes at least one selection marker. More preferably, it may be in the form of a phage, a plasmid, a

cosmid, a mini-chromosome, a virus, a retrovirus vector to which the polynucleotide is introduced.

As used herein, the term "transformant" refers to a host cell transformed with the vector, and means a transformant capable of producing a large amount of soluble, or soluble and crystalline NanR protein of the present invention, and also includes a transformant which is introduced with the NanR protein and thus is used to screen candidates for new drug development through NMR, etc, but is not limited thereto. The transformant introduced with the expression vector provided in the present invention may be, but is not particularly limited to, bacterial cells such as *E. coli*, *Streptomyces*, *Salmonella typhimurium*, etc.; yeast cells; fungus cells such as *Pichia pastoris*; insect cells such as *Drosophila*, *Spodoptera* Sf9 cell, etc.; animal cells such as CHO, COS, NSO, 293T, Bowes melanoma cells, etc.; or plants cells, which are transformed by introduction of the expression vector.

As used herein, the term "introduction" refers to delivery of the vector including the polynucleotide encoding the NanR protein into a host cell. This introduction may be performed by various methods known in the art, including calcium phosphate-DNA coprecipitation, DEAE dextran-mediated transfection, polybrene-mediated transfection, electroporation, microinjection, liposome fusion, lipofection and protoplast fusion. Also, term "transfection" means delivery of a desired material into a cell by means of infection using viral particles. In addition, the vector may be introduced into a host cell by gene bombardment. In the present invention, the introduction may be used interchangeably with transformation.

In still another aspect, the present invention provides a method for screening a substance regulating the interaction between NanR and the transcriptional control region of nan operon, which is a cluster of genes regulated by NanR, by utilizing three-dimensional structure of the complex of NanR protein and ManNAc-6P.

Preferably, the method may include the steps of (a) designing a tertiary structure of the complex using the atomic coordinates of the complex of NanR protein and ManNAc-6P, of which protein data bank accession code is 4IVN, that is, the atomic coordinates shown in Table 3; (b) preparing candidates binding to NanR using the tertiary structure thus designed; and (c) examining binding affinity of the candidate for NanR and its regulation of the interaction between NanR and the transcriptional control region of nan operon, but is not limited thereto.

Preferably, the method for screening a substance regulating the interaction between NanR and the transcriptional control region of nan operon may further include the step of determining the candidate as a growth inhibitor for bacteria having nan gene, if the corresponding candidate binding to NanR has the NanR binding affinity similar to or higher than that of ManNAc-6P and maintains or increases interaction between NanR and the transcriptional control region of nan operon, compared to a control group treated without the corresponding candidate. More preferably, the method may further include a method for screening a substance having a higher bacterial growth-inhibiting ability than the candidate screened in step (c) by using the designed tertiary structure.

Preferably, the method for screening a substance regulating interaction between NanR and the transcriptional control region of nan operon may further include the step of determining the candidate as a growth stimulant for bacteria having nan gene, if the corresponding candidate binding to NanR has the NanR binding affinity similar to or higher than that of ManNAc-6P and decreases interaction between

NanR and the transcriptional control region of nan operon, compared to a control group treated without the corresponding candidate. More preferably, the method may further include a method for screening a substance having a higher bacterial growth-stimulating ability than the candidate screened in step (c) by using the designed tertiary structure.

In the present invention, the tertiary structure of NanR protein/ManNAc-6P complex was demonstrated, and an electron density difference map showing the interaction of NanR protein and ManNAc-6P in the complex demonstrated that ManNAc-6P mainly binds to L9, L13, and L17 loop structures of the NanR LBD domain. Specifically, it was confirmed that Arg71 of a DNA binding domain (DBD) and Ala137, Ser138, His163, Ser182, Ser183, Ser184, Thr187, Glu229, Pro231, Gly234, or/and Lys240 of a ligand binding domain (LBD) of NanR protein and ManNAc-6P forms a hydrogen bond or a water-mediated hydrogen bond. Based on this information, therefore, a compound or a peptide capable of binding to NanR and regulating its activity can be designed or/and synthesized. The compound is preferably a small molecule compound, but is not limited thereto. In addition, because it is known that the ManNAc-6P binds to NanR to inhibit interaction between NanR and the transcriptional control region of nan operon, the NanR-binding peptides or compounds synthesized by using the NanR structural information may be substances that binds to NanR to inhibit its activity, like ManNAc-6P, but is not limited thereto. The substance may be a substance that maintains or stimulates the inhibitory effect of NanR on transcription of the nan gene through binding competition with ManNAc-6P, if it has the NanR binding affinity similar to or higher than that of ManNAc-6P but does not inhibit the activity.

In the present invention, the bacterial growth may occur in the presence of sialic acid, in particular, N-acetylneuraminic acid (Neu5Ac) as a carbon source.

NanR protein, ManNAc-6P, and the complex thereof are the same as described above. The atomic coordinates for the complex of NanR protein and ManNAc-6P can be obtained from the protein data bank (PDB) accession code or the atomic coordinates shown in Table 3. Atomic coordinates for the complex of NanR protein and ManNAc-6P and the complex of SeMet-NanR protein and ManNAc-6P are shown in Table 3.

Further, the atomic coordinates for the complex of NanR protein and ManNAc-6P and the complex of SeMet-NanR protein and ManNAc-6P can be stored in media for consecutive use in a calculating apparatus such as a computer. Typically, the coordinates can be stored in media (e.g., floppy disks, hard disks, compact disks, magneto-optical media, or electronic media) useful for storing large amounts of data, such as magnetic or optical media. Those skilled in the structural/computational chemistry are used to selecting the computer, storage media, networking and other device or technique.

Based on the three-dimensional structure of the complex of NanR protein and ManNAc-6P or the complex of SeMet-NanR protein and ManNAc-6P demonstrated in the present invention, information about various protein sites including binding sites can be provided by using a computer readable media containing the data of the atomic coordinates and/or three-dimensional structure. Through these procedures, reaction patterns of numerous drug candidates can be predicted without practical experiments, and only the resulting selected substances are subjected to experiments, leading to economic improvement in new drug development.

Further, step (a) of the screening method may include the steps of entering the data of the atomic coordinates for the

tertiary structure of the complex into a computer, together with a proper software program; and obtaining a three-dimensional protein structure for visualization and additional computer manipulation, but is not limited thereto.

If all or part of the tertiary structure of the NanR protein/ManNAc-6P complex or the SeMet-NanR protein/ManNAc-6P complex is used, a candidate regulating the binding of NanR protein and ManNAc-6P can be specifically selected or produced. As confirmed by the present inventors, binding of NanR and the transcriptional control region of nan operon is regulated by binding of NanR protein and ManNAc-6P, and thus a candidate regulating the binding of NanR protein and the transcriptional control region of nan operon can be screened by screening the candidate regulating the binding of NanR protein and ManNAc-6P.

The nan operon genes regulated by the transcriptional control region of nan operon are known to be involved in the Neu5Ac catabolism, that is, metabolism for utilizing Neu5Ac as a carbon source, and to play a very important role in proliferation and survival of bacteria in the host intestine in which Neu5Ac is abundant as a carbon source.

Therefore, the candidate which has the NanR binding affinity similar to or higher than that of ManNAc-6P and maintains or increases interaction between NanR and the transcriptional control region of nan operon, compared to a control group treated without the corresponding candidate, is able to prevent a reduction in the function of NanR as the nan operon transcriptional repressor due to binding with ManNAc-6P, and also to maintain or improve the function of NanR as the nan operon transcriptional repressor. Therefore, the candidate inhibits growth of the bacteria having nan operon genes, and shows the effects of reducing proliferation, survival and pathogenicity of the bacteria, and thus it can be determined as a bacterial growth inhibitor or an antibacterial agent.

Meanwhile, the candidate which has the NanR binding affinity similar to or higher than that of ManNAc-6P and decreases interaction between NanR and the transcriptional control region of nan operon, compared to a control group treated without the corresponding candidate, is able to inhibit the function of NanR as the nan operon transcriptional repressor instead of ManNAc-6P and to promote transcription of nan operon. Therefore, the candidate promotes growth of the bacteria having nan operon genes, and shows the effects of increasing proliferation, survival and pathogenicity of the bacteria, and thus it can be determined as a bacterial growth stimulant.

As used herein, the term "candidate" includes DNA, RNA, an antibody, a compound, a peptide or a complex thereof without limitation, as long as it is able to bind with NanR. That is, the candidate may include a substance which is predicted to have a structure capable of binding to the corresponding NanR site based on the analyzed tertiary structure of NanR/ManNAc-6P complex, or which is synthesized, prepared, or modified to have the structure capable of binding thereto, without limitation.

As used herein, the term "nan gene" means a series of genes which are needed for utilizing sialic acid as a carbon source and found in bacteria, and may be a cluster of genes required for catabolism of sialic acid, and in particular, may refer to genes in the form of an operon. Specifically, sialic acid may mean Neu5Ac, and expression of nan genes is up-regulated when sialic acid is supplied. nan genes play an important role in colonization and pathogenic activity of bacteria having nan genes. In the present invention, nan genes may be genes (nanA, nanTL, nanTS, nanTP, nanE or nanK genes, etc.) repressed by NanR protein, but are not

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limited thereto. For example, *E. coli* requires nanATEK operon for catabolism of Neu5Ac, and NanR is a repressor of this operon. The amino acid sequences and nucleotide sequences of the nan genes can be obtained from the known database such as GenBank at The National Center for Biotechnology Information, and genes represented by amino acid sequences of SEQ ID NOS. 3 to 8 or nucleotide sequences of SEQ ID NOS. 9 to 14 are preferred.

As used herein, the term "bacteria having nan genes" means bacteria that are able to utilize sialic acid as a carbon source because they have nan genes required for utilization of sialic acid as a carbon source. In the present invention, the bacteria may be *V. vulnificus*, *E. coli*, *Haemophilus influenzae*, or *V. cholerae*, but are not limited thereto, as long as they have nan genes. In particular, the bacteria having nan genes may be those having nan genes, of which transcription is repressed by NanR.

In still another aspect, the present invention provides a method for screening a substance regulating interaction between NanR and ManNAc-6P by using the three-dimensional structure of the complex of NanR protein and ManNAc-6P.

Specifically, the method may be a method for screening a substance regulating interaction between NanR and ManNAc-6P, including the steps of (a) designing a tertiary structure of the complex using the atomic coordinates of the complex of NanR protein and ManNAc-6P, of which protein data bank accession code is 4IVN, that is, the atomic coordinates shown in Table 3; (b) preparing candidates binding to NanR using the tertiary structure thus designed; and (c) examining whether the candidate regulates interaction between NanR protein and ManNAc-6P.

Herein, the terms and designing of the tertiary structure are the same as described above.

In step (c), if the candidate increases interaction between NanR and ManNAc-6P, compared to a control group treated without the corresponding candidate, the candidate is determined as a nan operon expression enhancer or as a bacterial growth stimulant. If the candidate decreases interaction between NanR and ManNAc-6P, compared to a control group treated without the corresponding candidate, the candidate is determined as a nan operon expression suppressor or as a bacterial growth inhibitor. Herein, regulation of the interaction between NanR and the transcriptional control region of nan operon can be also examined.

In still another aspect, the present invention provides an antibacterial composition including the bacterial growth inhibitor as screened above.

The screening method is the same as described above.

As used herein, the term "antibacterial composition" means a composition that functions to inhibit survival and/or growth of microorganisms. In particular, the composition may have an anti-bacterial activity against bacteria having nan genes, and specifically, an anti-bacterial activity against a microorganism selected from the group consisting of *V. vulnificus*, *E. coli*, *H. influenzae*, and *V. cholerae*, but is not limited thereto.

Further, the antibacterial composition of the present invention may be used for the prevention or treatment of various symptoms associated with bacteria having nan genes, in particular, septicemia.

As used herein, the term "septicemia" refers to a systemic inflammatory response caused by severe infection of microorganisms. Septicemia is a condition that is caused by the spread of microorganisms from one organ via the circulating blood, leading to systemic inflammatory response syndrome. However, systemic septicemia can be also caused by

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inflammatory response and production of inflammatory mediators in a part of the body, even though the microorganisms do not invade the bloodstream. In the present invention, septicemia may be caused by bacteria having nan genes, in particular, *V. vulnificus*, *E. coli*, *H. influenzae*, and *V. cholera*, but is not limited thereto.

The composition may further include a pharmaceutically acceptable carrier. As used herein, the term "pharmaceutically acceptable carrier" refers to a pharmaceutically acceptable material, composition or vehicle such as a liquid or solid filler, diluent, excipient, solvent or encapsulating material, which is involved in carrying or transporting any composition or component from one organ or a part of the body to another organ or a different region of the body. For administration, the composition of the present invention may include a pharmaceutically acceptable carrier, excipient, or diluent, in addition to the active ingredients described above. Examples of the carrier, excipient, and diluent may include lactose, dextrose, sucrose, sorbitol, mannitol, xylitol, erythritol, maltitol, starch, acacia rubber, alginate, gelatin, calcium phosphate, calcium silicate, cellulose, methyl cellulose, microcrystalline cellulose, polyvinylpyrrolidone, water, methylhydroxy benzoate, propylhydroxy benzoate, talc, magnesium stearate, and mineral oil.

According to the conventional methods, the composition of the present invention may be formulated into an oral preparation such as a powder, a granule, a tablet, a capsule, a suspension, an emulsion, a syrup, or an aerosol, an external preparation, suppository, or a sterilized injectable solution. In detail, such preparations may be prepared using diluents or excipients ordinarily employed, such as a filler, an extender, a binder, a wetting agent, a disintegrating agent, and a surfactant. Examples of the solid preparation for oral administration include a tablet, a pill, a powder, a granule, and a capsule, but are not limited thereto. The solid preparation may be prepared by mixing with at least one excipient such as starch, calcium carbonate, sucrose, lactose, or gelatin. Further, in addition to the excipients, lubricants such as magnesium stearate and talc may be used. The liquid formulation for oral administration may include a suspension, a liquid for internal use, an emulsion, a syrup or the like, but is not limited thereto. It may be prepared by adding various excipients such as a wetting agent, a sweetener, a flavor, or a preservative, in addition to general diluents such as water and liquid paraffin. Examples of the formulation for parenteral administration include an aseptic aqueous solution, a non-aqueous solvent, a suspension, an emulsion, a lyophilized agent, and suppository. As the non-aqueous solvent and suspension, propylene glycol, polyethylene glycol, plant oil such as olive oil, or an injectable ester such as ethyloleate may be used. As a suppository base, witepsol, macrogol, tween 61, cacao butter, lauric butter, glycerogelatin or the like may be used.

Hereinafter, the present invention will be described in more detail with reference to the following Examples. However, these Examples are for illustrative purposes only, and the invention is not intended to be limited by these Examples.

## Example 1

## Bacterial Strains, Plasmids, and Growth Conditions

The bacterial strains and plasmids used in the present invention are listed in the following Table 1.

TABLE 1

Strain or plasmid	Relevant characteristics <sup>a</sup>	Reference or source
<b>Bacterial strains</b>		
<i>V. vulnificus</i>		
MO6-24/O	Wild type; Clinical isolate; virulent	Wright AC, 1990
MORR	MO6-24/O with spontaneous Rif <sup>r</sup> mutation, virulent	Laboratory collection
MORSR	MO6-24/O with spontaneous Rif <sup>r</sup> , Sm <sup>r</sup> mutation, virulent	Laboratory collection
BS1209	MO6-24/O with nanR R57A <sup>b</sup>	This study
BS1210	MO6-24/O with nanR H163L <sup>b</sup>	This study
BS1213	MORSR with nanR H163L <sup>b</sup> ; Rif <sup>r</sup> , Sm <sup>r</sup>	This study
<i>E. coli</i>		
DH5 $\alpha$	supE44 ΔlacU169 ( $\phi$ 80 lacZ ΔM15) hsdR17 recA1 endA1 gyrA96 thi-1 relAI	Laboratory collection
SM10Δpir	thi thr leu tonA lacY supE recA::RP4-2-Tc::Mu Δ pir, Km <sup>r</sup> ; host for $\pi$ -requiring plasmids; conjugal donor	Miller VL, 1988
BW25113	lacI <sup>a</sup> rrmB <sub>T14</sub> ΔlacZ <sub>WJ16</sub> hsdR514 ΔaraBAD <sub>AH33</sub> ΔrrhaBAD <sub>LD78</sub>	Datesenko KA, 2000
BSE1201	BW25113 with ΔnanE	This study
BL21 (DE3)	F <sup>-</sup> ompT hsdS (r <sub>e</sub> <sup>-</sup> , m <sub>B</sub> <sup>-</sup> gal (DE3)	Laboratory collection
Plasmids		
pGEM-T easy	PCR product cloning vector; Ap <sup>r</sup>	Promega
pBS1201	pGEM-T easy with nanR gene orf; Ap <sup>r</sup>	This study
pDM4	Suicide vector; onR6K; Cm <sup>r</sup>	Milton DL, 1996
pBS1206	pDM4 with nanR R57A <sup>b</sup> ; Cm <sup>r</sup>	This study
pBS1209	pDM4 with nanR H163L <sup>b</sup> ; Cm <sup>r</sup>	This study
pNT-K20A	pGEM-T easy with nanR K20A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNT-K21A	pGEM-T easy with nanR K21A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNT-R23A	pGEM-T easy with nanR R23A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNT-R23L	pGEM-T easy with nanR R23L <sup>b</sup> ; Ap <sup>r</sup>	This study
pNT-R57A	pGEM-T easy with nanR R57A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNT-R57L	pGEM-T easy with nanR R57L <sup>b</sup> ; Ap <sup>r</sup>	This study
pNT-R60A	pGEM-T easy with nanR R60A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNT-R60L	pGEM-T easy with nanR R60L <sup>b</sup> ; Ap <sup>r</sup>	This study
pNT-K65A	pGEM-T easy with nanR K65A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNT-S138A	pGEM-T easy with nanR S138A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNT-H163A	pGEM-T easy with nanR H163A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNT-H163L	pGEM-T easy with nanR H163L <sup>b</sup> ; Ap <sup>r</sup>	This study
pNT-S182A	pGEM-T easy with nanR S182A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNT-S184A	pGEM-T easy with nanR S184A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNT-T187A	pGEM-T easy with nanR T187A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNT-E229L	pGEM-T easy with nanR E229L <sup>b</sup> ; Ap <sup>r</sup>	This study
pNT-K240A	pGEM-T easy with nanR K240A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNT-K240M	pGEM-T easy with nanR K240M <sup>b</sup> ; Ap <sup>r</sup>	This study
pBAD-24BS	pBAD24 with unique XbaI site instead of XbaI site; Ap <sup>r</sup>	Kim Y, 2010
pNB-WT	pBAD-24BS with nanR; Ap <sup>r</sup>	This study
pNB-K20A	pBAD-24BS with nanR K20A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNB-K21A	pBAD-24BS with nanR K21A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNB-R23A	pBAD-24BS with nanR R23A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNB-R23L	pBAD-24BS with nanR R23L <sup>b</sup> ; Ap <sup>r</sup>	This study
pNB-R57A	pBAD-24BS with nanR R57A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNB-R57L	pBAD-24BS with nanR R57L <sup>b</sup> ; Ap <sup>r</sup>	This study
pNB-R60A	pBAD-24BS with nanR R60A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNB-R60L	pBAD-24BS with nanR R60L <sup>b</sup> ; Ap <sup>r</sup>	This study
pNB-K65A	pBAD-24BS with nanR K65A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNB-S138A	pBAD-24BS with nanR S138A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNB-H163A	pBAD-24BS with nanR H163A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNB-H163L	pBAD-24BS with nanR H163L <sup>b</sup> ; Ap <sup>r</sup>	This study
pNB-S182A	pBAD-24BS with nanR S182A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNB-S184A	pBAD-24BS with nanR S184A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNB-T187A	pBAD-24BS with nanR T187A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNB-E229L	pBAD-24BS with nanR E229L <sup>b</sup> ; Ap <sup>r</sup>	This study
pNB-K240A	pBAD-24BS with nanR K240A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNB-K240M	pBAD-24BS with nanR K240M <sup>b</sup> ; Ap <sup>r</sup>	This study
pBBR_lux	Broad host range vector containing luxCDABE operon; Cm <sup>r</sup>	Lenz DH, 2004
pBS0915	pBBR_lux with P <sub>nanT<sub>p</sub></sub> ; Cm <sup>r</sup>	This study
pHis-Parallel1	Protein expression vector; Ap <sup>r</sup>	Sheffield P, 1999
pBS0820	pHIS-Parallel1 with nanR; Ap <sup>r</sup>	Kim BS, 2011
pNH-R57A	pHis-Parallel1 with nanR R57A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNH-R60A	pHis-Parallel1 with nanR R60A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNH-S138A	pHis-Parallel1 with nanR S138A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNH-H163L	pHis-Parallel1 with nanR H163L <sup>b</sup> ; Ap <sup>r</sup>	This study
pNH-S182A	pHis-Parallel1 with nanR S182A <sup>b</sup> ; Ap <sup>r</sup>	This study
pNH-S184A	pHis-Parallel1 with nanR S184A <sup>b</sup> ; Ap <sup>r</sup>	This study

TABLE 1-continued

Strain or plasmid	Relevant characteristics <sup>a</sup>	Reference or source
pNH-E229L	pHis-Parallel1 with nanR E229L <sup>b</sup> ; Ap <sup>r</sup>	This study
pNH-K240A	pHis-Parallel1 with nanR K240A <sup>b</sup> ; Ap <sup>r</sup>	This study
pKD46	P <sub>BAD</sub> -gamt-beta-exo oriR101 repA101 <sup>c</sup> ; Ap <sup>r</sup>	Datesenko KA, 2000
pKD13	FRT Km <sup>r</sup> FRT PS1 PS4 oriR6K <sup>c</sup> ; Ap <sup>r</sup>	Datesenko KA, 2000
pCP20	c/875 ΔP <sub>αfp</sub> oriP SC101 <sup>c</sup> ; Ap <sup>r</sup> , Cm <sup>r</sup>	Datesenko KA, 2000
pBS0921	pRLG770 with P <sub>nanE</sub> ; Ap <sup>r</sup>	Kim BS, 2011

<sup>a</sup>Ap<sup>r</sup> represents ampicillin resistant; Cm<sup>r</sup> represents Chloramphenicol resistant; Km<sup>r</sup> represents kanamycin resistant; Rif<sup>r</sup> represents rifampicin resistant; Sm<sup>r</sup> represents streptomycin resistant.

<sup>b</sup>In the representation of mutant, first letter indicates an original amino acid, number indicates the position of mutation, and the last letter indicates an amino acid substituted due to mutation.

Unless noted otherwise, the *E. coli* and *V. vulnificus* strains were grown at 37° C. in LB (Luria-Bertani) medium and at 30° C. in LB supplemented with 2% NaCl, respectively. Where appropriate, antibiotics were added to medium at the following concentrations: 10 µg/ml chloramphenicol, 100 µg/ml ampicillin, and 100 µg/ml kanamycin for *E. coli*, and 3 µg/ml chloramphenicol, 100 µg/ml rifampicin, and 100 µg/ml streptomycin for *V. vulnificus*. M9 minimal medium was supplemented with the appropriate carbon sources (5 mM Neu5Ac alone, or 5 mM Neu5Ac, 10 mM D-xylose, and 10 mM L-proline). ManNAc-6P was purchased from Carbosynth (Berkshire, UK). All other chemicals were purchased from Sigma (St. Louis, Mo.).

#### Example 2

##### Construction of Plasmids and Strains

The nanR gene was amplified by PCR and cloned into the pGEM-T easy vector to form the pBS1201 construct. Site-directed mutations were introduced into this plasmid using the QuickChange™ Site-Directed Mutagenesis Kit (Agilent). The WT or mutant nanR genes were then subcloned into the NcoI and Xhol sites of the pBAD-24BS or pHis-parallel expression vector to construct the pNB- and pNH-plasmids, respectively. The R57A and H163L mutant nanR genes were subcloned into the SphI and SpeI sites of pDM4 to form pBS1208 and 1209, respectively.

To generate the NanR-dependent luciferase promoter-reporter plasmid (pBS0915), the intergenic region between nanE and nanTP was liberated from the pBS0909 plasmid and ligated with BamHI-digested pBBR\_lux. To construct the nanR R57A mutant *V. vulnificus* strain (BS1209) by homologous recombination, *E. coli* SM10λ pir,tra (containing pBS1208) (Miller and Mekalanos, J Bacteriol 170(6): pp 2575-2583, 1988) was used as a conjugal donor to MO6-24/O. Similarly, *E. coli* SM10λ pir,tra containing pBS1209 was used as a conjugal donor in conjunction with either MO6-24/O or MORSR to generate the nanR H163L mutants (BS1210 or BS1213, as indicated in Table 1). Conjugation and isolation of the transconjugants were performed as described previously. To construct the nanE deletion mutant *E. coli* strain (BSE1201), the lambda Red-recombineering method was used, as known previously. Briefly, the kanamycin resistant (KmR) cassette from pKD13 was PCR amplified and then electroporated into the BW25113 strain containing pKD46. Insertion of the KmR cassette into nanE was confirmed by PCR, and the cassette was subsequently removed from the chromosome by transforming pCP20 into kanamycin resistant cells. After verifying the deletion of nanE by PCR, the cells were maintained at 37° C. for the plasmid curing.

#### Example 3

##### *E. coli* Dual Plasmid System

*E. coli* strains were co-transformed with a luciferase reporter plasmid (pBS0915) and one of the pNB-series of plasmids that express NanR. Cells were cultured overnight and then diluted into the appropriate fresh media (supplemented M9 in FIG. 3; LB in FIG. 4) containing arabinose (0.002%), and incubated at 37° C. until cells grew to early exponential phase. Relative luminescence unit (RLU) was calculated by dividing the luminescence by the A<sub>600</sub>, as described previously. For screening ligand-sensing residues, the BSE1201 strain (ΔaraBAD ΔnanE) was used as the host cell instead of DH5α to ensure that the arabinose was not used as a carbon source and that ManNAc-6P generated from Neu5Ac was accumulated in the cell.

#### Example 4

##### In Vitro Transcription Assay and qRT-PCR

In vitro transcription assays with wild-type or mutant NanR proteins were performed according to procedures described previously. RNA extraction, cDNA synthesis, and real time PCR amplification of the cDNA were performed as described previously.

#### Example 5

##### In Vitro Growth Defect and Mouse Experiments

MO6-24/O (WT), BS1209 (R57A mutant), and BS1210 (H163L mutant) strains were cultured overnight and then serially diluted in PBS. 10 mL of each dilution were spotted onto M9 minimal media supplemented with 5 mM Neu5Ac only or 5 mM Neu5Ac, 10 mM D-xylose, and 10 mM L-proline. The growth and phenotype of the strains were examined after incubating at 30° C. for 24 hours.

For the mouse intestine colonization competition assay, 10 mice (6 weeks old, female ICR) were provided with drinking water containing rifampicin (50 µg/ml) for 24 hours to eliminate resident bacteria. After a starvation period without food and water, mice were intragastrically injected with the bacterial mixture of MORR (WT; RifR) and BS1213 (H163L mutant; RifR, SmR) (approximately 1x10<sup>6</sup> CFU per strain). At 12 hours after infection, mice were euthanized and the small intestines were collected and homogenized in 5 ml of PBS. Equal amounts of neat or diluted homogenates were spread onto LBS agar supplemented with 2% NaCl and either rifampicin alone (to count the sum of WT and H163L mutant cells), or rifampicin and streptomycin (to count the H163L mutant cells only). The

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competitive index was calculated by dividing the recovered mutant/WT ratio by the inoculated mutant/WT ratio. For the mouse survival test, five mice per group were intragastrically infected with  $4 \times 10^8$  CFU of either MO6-24/O (WT) or BS1210 (H163L mutant) strains and monitored for 1 day.

All animal experiments were performed according to the recommended procedures for the care and use of laboratory animals from the Institute of Laboratory Animal Resource at Seoul National University. The protocol was approved by the Committee on the Ethics of Animal Experiments of Seoul National University (Institutional Animal Care and Use Committee approval number: SNU-111130-2).

#### Example 6

##### Western Blot, EMSA, and DNaseI Footprinting Assay

Purified His-NanR protein was used to raise a primary polyclonal antibody by immunizing European rabbits (*Oryctolagus cuniculus*). First, after one primary injection containing 500 µg of protein, three boosters containing 200 µg of protein was performed at 2-week intervals. Western blotting, EMSA and DNaseI footprinting assays were performed according to typical procedures known in the art.

#### Example 7

##### Protein Expression and Purification

Expression and purification of the NanR protein was performed according to typical procedures known in the art. To obtain the selenomethionine (SeMet)-substituted NanR protein, the methionine auxotroph *E. coli* B834 (DE3) strain (Novagen) was grown in minimal medium supplemented with 50 mg/ml SeMet. The SeMet-substituted NanR was purified and obtained from the culture by the addition of 5 mM methionine to all buffers.

#### Example 8

##### Crystallization, Diffraction, and Structure Determination

Crystallization trials of the purified NanR protein performed using the sitting drop vapor-diffusion method at 21°

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C. were unsuccessful. However, the crystals were obtained when NanR and ManNAc-6P were mixed in a 1:100 molar ratio and incubated on ice for 2 hours.

The crystallization of complex was optimized under the following conditions: 10% PEG 2,000 MME, 0.1 M ammonium sulfate, 0.3 M sodium formate, 3% PGA-LM, and 0.1 M sodium acetate (pH 5.0 to 5.5). Crystals appeared within a day and were grown for a further 5 days for diffraction experiments. The complex crystals were transferred to a cryoprotectant solution containing 10% PEG 2,000 MME, 0.1 M ammonium sulfate, 0.3 M sodium formate, 3% PGA-LM, 0.1 M sodium acetate (pH 5.5), and 30% glycerol, and then placed immediately in a -173° C. nitrogen gas stream.

Diffraction data for the complex crystals were collected at 1.9 Å resolution. SeMet-substituted complex crystals were grown under the same crystallization conditions as described above. Single wavelength anomalous diffraction data for the SeMet-substituted crystals were collected at 2.4 Å resolution. All data were processed with HKL2000 package. The structure of the NanR/ManNAc-6P complex was determined by analyzing the anomalous signals from Se atoms with the SOLVE program. Density modification and subsequent automated model building were performed using the RESOLVE program. The complex crystal structure was solved at 1.9 Å resolution by molecular replacement with the MOLREP program using the partially refined model of the SeMet crystal. The complex crystal structure was revised using the COOT program and refined using the REFMAC5. The refinement included the translation-liberation-screw (TLS) procedure. The final refined model resulted in  $R_{free}$  and  $R_{cryst}$  values of 0.235 and 0.183, respectively. No density was visible for the Met1 to Lys5 and Glu82 to Glu90 regions of NanR-A, and the Met1 to Lys5, Thr81 to Gly91, and Asn278 regions of NanR-B. These residues were not included in the model. The model contained 525 amino acid residues, two ManNAc-6P molecules, and 221 water molecules, and satisfied the quality criteria limits of the PROCHECK program.

The crystallographic data statistics are summarized in the following Table 2.

TABLE 2

Data collection and refinement statistics for ManNAc-6P/NanR complex		
Dataset	SeMet-NanR complexed with ManNAc-6P	NanR complexed with ManNAc-6P
Wavelength	0.97917	1.0000
Space group	P3 <sub>1</sub> 21	P3 <sub>1</sub> 21
Unit cell (Å)	$a = b = 109.84, c = 83.38$ $\alpha = \beta = 90^\circ, \gamma = 120^\circ$	$a = b = 109.21, c = 82.47$ $\alpha = \beta = 90^\circ, \gamma = 120^\circ$
Resolution, (Å)	50.0-2.40 (2.49-2.40)	50.0-1.90 (1.93-1.90)
No. of total reflections	508,845	330,826
No. of unique reflections	23,076	45,033
Redundancy	22.1 (22.47)	7.3 (7.3)
Completeness (%)	99.9 (100.0)	99.9 (100.0)
$R_{sym}$ (%) <sup>a</sup>	9.4 (31.5)	4.7 (47.3)
I/σ(I)	50.27 (12.28)	42.05 (3.15)
Refinement		
Resolution (Å)		30.0-1.90
Reflections in work/test sets		42,737/2,268
$R_{work}/R_{free}$ (%) <sup>b,c</sup>		18.3/23.5

TABLE 2-continued

Data collection and refinement statistics for ManNAc-6P/NanR complex		
Dataset	SeMet-NanR complexed with ManNAc-6P	NanR complexed with ManNAc-6P
<u>R.m.s. deviations</u>		
Bond lengths (Å)		0.021
Bond angles (°)		2.126
Model composition	525 residues 221 waters 2 ManNAc-6P	
<u>Geometry</u>		
Favored regions (%)		98.8
Allowed regions (%)		1.2
PDB accession code	4IVN	

<sup>a</sup>R<sub>sym</sub> =  $\sum |I_i - \langle I \rangle| / \sum |I|$  where I<sub>i</sub> is the intensity of the i-th observation and <I> is the mean intensity of the reflections.

<sup>b</sup>R<sub>wrk</sub> =  $\sum |F_{obs} - |F_{calc}|| / \sum |F_{obs}|$  where F<sub>calc</sub> and F<sub>obs</sub> are measured and calculated, respectively.

<sup>c</sup>R<sub>free</sub> =  $\sum |F_{obs} - |F_{calc}|| / \sum |F_{obs}|$  where all reflections belong to a test set of randomly selected data.

TABLE 3

---

HEADER TRANSCRIPTION REGULATOR 23-JAN-13 4IVN

TITLE THE VIBRIO VULNIFICUS NANR PROTEIN COMPLEXED WITH MANNAC-6P

COMPND MOL\_ID: 1:

COMPND 2 MOLECULE: TRANSCRIPTIONAL REGULATOR :

COMPND 3 CHAIN: A. B:

COMPND 4 SYNONYM: NANR:

COMPND 5 ENGINEERED: YES

SOURCE MOL\_ID: 1:

SOURCE 2 ORGANISM\_SCIENTIFIC: VIBRIO VULNIFICUS:

SOURCE 3 ORGANISM\_TAXID: 196600:

SOURCE 4 STRAIN: YJ016:

SOURCE 5 EXPRESSION\_SYSTEM: ESCHERICHIA COLI:

SOURCE 6 EXPRESSION\_SYSTEM\_TAXID: 562

KEYWDS ISOMERASE FOLD. NAN OPERON REGULATOR FOR SIALIC ACID CATABOLISM.

KEYWDS 2 TRANSCRIPTION REGULATOR

EXPDTA X-RAY DIFFRACTION

AUTHOR J. HWANG, M. H. KIM

REVDAT 2 20-NOV-13 4INV: JRNL

REVDAT 1 17-JUL-13 4IVN 0

JRNL AUTH J. HWANG, B. S. KIM, S. Y. JANG, J. G. LIM, D. J. YOU, H. S. JUNG, T. K. OH,

JRNL AUTH 2 J. O. LEU, S. H. CHOI, M. H. KIM

JRNL TITL STRUCTURAL INSIGHTS INTO THE REGULATION OF SIALIC ACID

JRNL TITL 2 CATABOLISM BY THE VIBRIO VULNIFICUS TRANSCRIPTIONAL

JRNL TITL 3 REPRESSOR NANR

JRNL REF PROC. NATL. ACAD. SCI. USA V. 110 E2829 2013

JRNL REFN ISSN 0027-8424

TABLE 3-continued

JRNL PMID 23832782

JRNL DOI 10.1073/PNAS.1302859110

REMARK 2

REMARK 2 RESOLUTION. 1.90 ANGSTROMS.

REMARK 3

REMARK 3 REFINEMENT.

REMARK 3 PROGRAM: REFMAC 5.6.0117

REMARK 3 AUTHORS: MURSHUDOV. VAGIN. DODSON

REMARK 3

REMARK 3 REFINEMENT TARGET: MAXIMUM LIKELIHOOD

REMARK 3

REMARK 3 DATA USED IN REFINEMENT.

REMARK 3 RESOLUTION RANGE HIGH (ANGSTROMS): 1.90

REMARK 3 RESOLUTION RANGE LOW (ANGSTROMS): 30.00

REMARK 3 DATA CUTOFF (SIGMA(F)): NULL

REMARK 3 COMPLETENESS FOR RANGE (%): 99.9

REMARK 3 NUMBER OF REFLECTIONS: 42737

REMARK 3

REMARK 3 FIT TO DATA USED IN REFINEMENT.

REMARK 3 CROSS-VALIDATION METHOD: THROUGHOUT

REMARK 3 FREE R VALUE TEST SET SELECTION: RANDOM

REMARK 3 R VALUE (WORKING + TEST SET): 0.183

REMARK 3 R VALUE (WORKING SET): 0.181

REMARK 3 FREE R VALUE: 0.235

REMARK 3 FREE R VALUE TEST SET SIZE (%): 5.000

REMARK 3 FREE R VALUE TEST SET COUNT: 2268

REMARK 3

REMARK 3 FIT IN THE HIGHEST RESOLUTION BIN.

REMARK 3 TOTAL NUMBER OF BINS USED: 20

REMARK 3 BIN RESOLUTION RANGE HIGH (A): 1.90

REMARK 3 BIN RESOLUTION RANGE LOW (A): 1.95

REMARK 3 REFLECTION IN BIN (WORKING SET): 2947

REMARK 3 BIN COMPLETENESS (WORKING + TEST) (%): 99.97

REMARK 3 BIN R VALUE (WORKING SET): 0.2690

REMARK 3 BIN FREE R VALUE SET COUNT: 162

REMARK 3 BIN FREE R VALUE: 0.3630

REMARK 3

REMARK 3 NUMBER OF NON-HYDROGEN ATOMS USED IN REFINEMENT.

REMARK 3 PROTEIN ATOMS: 4004

REMARK 3 NUCLEIC ACID ATOMS: 0

TABLE 3-continued

REMARK 3 HETEROGEN ATOMS: 38

REMARK 3 SOLVENT ATOMS: 221

REMARK 3

REMARK 3 B VALUES.

REMARK 3 FROM WILSON PLOT (A\*\*2): NULL

REMARK 3 MEAN B VALUE (OVERALL. A\*\*2): 42.47

REMARK 3 OVERALL ANISOTROPIC B VALUE.

REMARK 3 B11 (A\*\*2): -2.27000

REMARK 3 B22 (A\*\*2): -2.27000

REMARK 3 B33 (A\*\*2): 3.41000

REMARK 3 B12 (A\*\*2): -1.14000

REMARK 3 B13 (A\*\*2): 0.00000

REMARK 3 B23 (A\*\*2): 0.00000

REMARK 3

REMARK 3 ESTIMATED OVERALL COORDINATE ERROR.

REMARK 3 ESU BASED ON R VALUE (A): 0.142

REMARK 3 ESU BASED ON FREE R VALUE (A): 0.144

REMARK 3 ESU BASED ON MAXIMUM LIKELIHOOD (A): 0.106

REMARK 3 ESU FOR B VALUES BASED ON MAXIMUM LIKELIHOOD (A\*\*2): 7.038

REMARK 3

REMARK 3 CORRELATION COEFFICIENTS.

REMARK 3 CORRELATION COEFFICIENT FO-FC: 0.956

REMARK 3 CORRELATION COEFFICIENT FO-FC FREE: 0.941

REMARK 3

REMARK 3 RMS DEVIATIONS FROM IDEAL VALUES COUNT RMS WEIGHT

REMARK 3 BOND LENGTHS REFINED ATOMS (A): 4096 : 0.021 : 0.019

REMARK 3 BOND LENGTHS OTHERS (A): NULL : NULL : NULL

REMARK 3 BOND ANGLES REFINED ATOMS (DEGREES): 5548 : 2.126 : 1.982

REMARK 3 BOND ANGLES OTHERS (DEGREES): NULL : NULL : NULL

REMARK 3 TORSION ANGLES, PERIOD 1 (DEGREES): 521 : 6.387 : 5.000

REMARK 3 TORSION ANGLES, PERIOD 2 (DEGREES): 165 : 35.450 : 23.576

REMARK 3 TORSION ANGLES, PERIOD 3 (DEGREES): 716 : 16.845 : 15.000

REMARK 3 TORSION ANGLES, PERIOD 4 (DEGREES): 32 : 18.969 : 15.000

REMARK 3 CHIRAL-CENTER RESTRAINTS (A\*\*3): 669 : 0.146 : 0.200

REMARK 3 GENERAL PLANES REFINED ATOMS (A): 2984 : 0.010 : 0.021

REMARK 3 GENERAL PLANES OTHERS (A): NULL : NULL : NULL

REMARK 3 NON-BONDED CONTACTS REFINED ATOMS (A): NULL : NULL : NULL

REMARK 3 NON-BONDED CONTACTS OTHERS (A): NULL : NULL : NULL

REMARK 3 NON-BONDED TORSION REFINED ATOMS (A): NULL : NULL : NULL

REMARK 3 NON-BONDED TORSION OTHERS (A): NULL : NULL : NULL

TABLE 3-continued

REMARK 3 H-BOND (X...Y) REFINED ATOMS (A): NULL : NULL : NULL

REMARK 3 H-BOND (X...Y) OTHERS (A): NULL : NULL : NULL

REMARK 3 POTENTIAL METAL-ION REFINED ATOMS (A): NULL : NULL : NULL

REMARK 3 POTENTIAL METAL-ION OTHERS (A): NULL : NULL : NULL

REMARK 3 SYMMETRY VDW REFINED ATOMS (A): NULL : NULL : NULL

REMARK 3 SYMMETRY VDW OTHERS (A): NULL : NULL : NULL

REMARK 3 SYMMETRY H-BOND REFINED ATOMS (A): NULL : NULL : NULL

REMARK 3 SYMMETRY H-BOND OTHERS (A): NULL : NULL : NULL

REMARK 3 SYMMETRY METAL-ION REFINED ATOMS (A): NULL : NULL : NULL

REMARK 3 SYMMETRY METAL-ION OTHERS (A): NULL : NULL : NULL

REMARK 3

REMARK 3 ISOTROPIC THERMAL FACTOR RESTRAINTS. COUNT RMS WEIGHT

REMARK 3 MAIN-CHAIN BOND REFINED ATOMS (A\*\*2): NULL : NULL : NULL

REMARK 3 MAIN-CHAIN BOND OTHER ATOMS (A\*\*2): NULL : NULL : NULL

REMARK 3 MAIN-CHAIN ANGLE REFINED ATOMS (A\*\*2): NULL : NULL : NULL

REMARK 3 MAIN-CHAIN ANGLE OTHER ATOMS (A\*\*2): NULL : NULL : NULL

REMARK 3 SIDE-CHAIN BOND REFINED ATOMS (A\*\*2): NULL : NULL : NULL

REMARK 3 SIDE-CHAIN BOND OTHER ATOMS (A\*\*2): NULL : NULL : NULL

REMARK 3 SIDE-CHAIN ANGLE REFINED ATOMS (A\*\*2): NULL : NULL : NULL

REMARK 3 SIDE-CHAIN ANGLE OTHER ATOMS (A\*\*2): NULL : NULL : NULL

REMARK 3 LONG RANGE B REFINED ATOMS (A\*\*2): NULL : NULL : NULL

REMARK 3 LONG RANGE B OTHER ATOMS (A\*\*2): NULL : NULL : NULL

REMARK 3

REMARK 3 ANISOTROPIC THERMAL FACTOR RESTRAINTS. COUNT RMS WEIGHT

REMARK 3 RIGID-BOND RESTRAINTS (A\*\*2): NULL : NULL : NULL

REMARK 3 SPHERICITY: FREE ATOMS (A\*\*2): NULL : NULL : NULL

REMARK 3 SPHERICITY: BONDED ATOMS (A\*\*2): NULL : NULL : NULL

REMARK 3

REMARK 3 NCS RESTRAINTS STATISTICS

REMARK 3 NCS TYPE: LOCAL

REMARK 3 NUMBER OF DIFFERENT NCS PAIRS: 1

REMARK 3 GROUP CHAIN1 RANGE CHAIN2 RANGE COUNT RMS WEIGHT

REMARK 3 1 A 6 277 B 6 277 366 0.15 0.05

REMARK 3

REMARK 3 TLS DETAILS

REMARK 3 NUMBER OF TLS GROUPS: 4

REMARK 3

REMARK 3 TLS GROUP: 1

REMARK 3 NUMBER OF COMPONENTS GROUP: 1

REMARK 3 COMPONENTS C SSSEQI TO C SSSEQI

TABLE 3-continued

REMARK 3 RESIDUE RANGE: A 5 A 81

REMARK 3 ORIGIN FOR THE GROUP (A): 17.7780 -5.8420 8.8420

REMARK 3 T TENSOR

REMARK 3 T11: 0.0449 T22: 0.1497

REMARK 3 T33: 0.1331 T12: 0.0044

REMARK 3 T13: -0.0544 T23: -0.0500

REMARK 3 L TENSOR

REMARK 3 L11: 3.4953 L22: 2.2054

REMARK 3 L33: 2.1117 L12: -0.5910

REMARK 3 L13: 0.7559 L23: 0.5527

REMARK 3 S TENSOR

REMARK 3 S11: 0.2213 S12: 0.2193 S13: -0.1430

REMARK 3 S21: -0.1435 S22: -0.2725 S23: 0.3625

REMARK 3 S31: 0.1735 S32: -0.1381 S33: 0.0512

REMARK 3

REMARK 3 TLS GROUP: 2

REMARK 3 NUMBER OF COMPONENTS GROUP: 1

REMARK 3 COMPONENTS C SSSEQI TO C SSSEQI

REMARK 3 RESIDUE RANGE: A 91 A 278

REMARK 3 ORIGIN FOR THE GROUP (A): 45.2910 8.9760 13.2780

REMARK 3 T TENSOR

REMARK 3 T11: 0.0570 T22: 0.0852

REMARK 3 T33: 0.0778 T12: 0.0414

REMARK 3 T13: 0.0370 T23: 0.0537

REMARK 3 L TENSOR

REMARK 3 L11: 0.5889 L22: 0.6680

REMARK 3 L33: 0.8170 L12: -0.0380

REMARK 3 L13: 0.1453 L23: 0.1381

REMARK 3 S TENSOR

REMARK 3 S11: 0.0958 S12: 0.1783 S13: 0.0946

REMARK 3 S21: -0.1548 S22: -0.0145 S23: -0.0391

REMARK 3 S31: -0.0769 S32: -0.1389 S33: -0.0812

REMARK 3

REMARK 3 TLS GROUP: 3

REMARK 3 NUMBER OF COMPONENTS GROUP: 1

REMARK 3 COMPONENTS C SSSEQI TO C SSSEQI

REMARK 3 RESIDUE RANGE: B 6 B 80

REMARK 3 ORIGIN FOR THE GROUP (A): 82.6480 3.2410 8.3210

REMARK 3 T TENSOR

REMARK 3 T11: 0.0362 T22: 0.1890

TABLE 3-continued

REMARK 3 T33: 0.2221 T12: 0.0038

REMARK 3 T13: 0.0694 T23: -0.0873

REMARK 3 L TENSOR

REMARK 3 L11: 1.7284 L22: 1.3274

REMARK 3 L33: 2.4459 L12: 0.2465

REMARK 3 L13: -1.0201 L23: 0.0720

REMARK 3 S TENSOR

REMARK 3 S11: 0.0630 S12: 0.0848 S13: -0.0810

REMARK 3 S21: -0.1023 S22: 0.0872 S23: -0.1534

REMARK 3 S31: -0.0463 S32: 0.4472 S33: -0.1502

REMARK 3

REMARK 3 TLS GROUP: 4

REMARK 3 NUMBER OF COMPONENTS GROUP: 1

REMARK 3 COMPONENTS C SSSEQI TO C SSSEQI

REMARK 3 RESIDUE RANGE: B 92 B 277

REMARK 3 ORIGIN FOR THE GROUP (A): 55.3100 -10.8280 14.5630

REMARK 3 T TENSOR

REMARK 3 T11: 0.0555 T22: 0.0233

REMARK 3 T33: 0.1270 T12: 0.0218

REMARK 3 T13: -0.0027 T23: -0.0342

REMARK 3 L TENSOR

REMARK 3 L11: 0.9177 L22: 0.6147

REMARK 3 L33: 0.8648 L12: -0.0846

REMARK 3 L13: -0.0381 L23: 0.0605

REMARK 3 S TENSOR

REMARK 3 S11: 0.1126 S12: 0.1400 S13: -0.1727

REMARK 3 S21: -0.1213 S22: -0.0199 S23: -0.1418

REMARK 3 S31: 0.1198 S32: 0.0177 S33: -0.0927

REMARK 3

REMARK 3 BULK SOLVENT MODELLING

REMARK 3 METHOD USED: MASK

REMARK 3 PARAMETERS FOR MASK CALCULATION

REMARK 3 VDW PROBE RADIUS: 1.20

REMARK 3 ION PROBE RADIUS: 0.80

REMARK 3 SHRINKAGE RADIUS: 0.80

REMARK 3

REMARK 3 OTHER REFINEMENT REMARKS: NULL

REMARK 4

REMARK 4 4IVN COMPLIES WITH FORMAT V. 3.30. 13-JUL-11

REMARK 100

## TABLE 3-continued

REMARK 100 THIS ENTRY HAS BEEN PROCESSED BY PDBJ ON 28-JAN-13.

REMARK 100 THE RCSB ID CODE IS RCSB077266.

REMARK 200

REMARK 200 EXPERIMENTAL DETAILS

REMARK 200 EXPERIMENT TYPE: X-RAY DIFFRACTION

REMARK 200 DATE OF DATA COLLECTION: 08-JUN-12

REMARK 200 TEMPERATURE (KELVIN): 100.15

REMARK 200 PH: 5.0

REMARK 200 NUMBER OF CRYSTALS USED: 1

REMARK 200

REMARK 200 SYNCHROTRON (Y/N): Y

REMARK 200 RADIATION SOURCE: PHOTON FACTORY

REMARK 200 BEAMLINE: BL -17A

REMARK 200 X-RAY GENERATOR MODEL: NULL

REMARK 200 MONOCHROMATIC OR LAUE (M/L): M

REMARK 200 WAVELENGTH OR RANGE (Å): 1.0000

REMARK 200 MONOCHROMATOR: NULL

REMARK 200 OPTICS: NULL

REMARK 200

REMARK 200 DETECTOR TYPE: CCD

REMARK 200 DETECTOR MANUFACTURER: NULL

REMARK 200 INTENSITY-INTEGRATION SOFTWARE: NULL

REMARK 200 DATA SCALING SOFTWARE: NULL

REMARK 200

REMARK 200 NUMBER OF UNIQUE REFLECTIONS: 45033

REMARK 200 RESOLUTION RANGE HIGH (Å): 1.900

REMARK 200 RESOLUTION RANGE LOW (Å): 50.000

REMARK 200 REJECTION CRITERIA (SIGMA00): 2.000

REMARK 200

REMARK 200 OVERALL.

REMARK 200 COMPLETENESS FOR RANGE (%): 99.9

REMARK 200 DATA REDUNDANCY: NULL

REMARK 200 R MERGE (1): NULL

REMARK 200 R SYM (1): NULL

REMARK 200 <I/SIGMA(1)> FOR THE DATA SET: NULL

REMARK 200

REMARK 200 IN THE HIGHEST RESOLUTION SHELL

REMARK 200 HIGHEST RESOLUTION SHELL. RANGE HIGH (Å): 1.90

REMARK 200 HIGHEST RESOLUTION SHELL. RANGE LOW (Å): 1.93

REMARK 200 COMPLETENESS FOR SHELL (%): 100.0

---

REMARK 200 DATA REDUNDANCY IN SHELL: NULL

REMARK 200 R MERGE FOR SHELL (1): NULL

REMARK 200 R SYM FOR SHELL (1): NULL

REMARK 200 <I/SIGMA(1)> FOR SHELL: NULL

REMARK 200

REMARK 200 DIFFRACTION PROTOCOL: SINGLE WAVELENGTH

REMARK 200 METHOD USED TO DETERMINE THE STRUCTURE: MOLECULAR REPLACEMENT

REMARK 200 SOFTWARE USED: NULL

REMARK 200 STARTING MODEL: NULL

REMARK 200

REMARK 200 REMARK: NULL

REMARK 280

REMARK 280 CRYSTAL

REMARK 280 SOLVENT CONTENT. VS (%): 47.42

REMARK 280 MATTHEWS COEFFICIENT. VM (ANGSTROMS\*\*3/DA): 2.34

REMARK 280

REMARK 280 CRYSTALLIZATION CONDITIONS: 10% PEG 2000 MME. 0.1M AMMONIUM

REMARK 280 SULFATE. 0.3M SODIUM FORMATE. 3% PGA-LM. 0.1M SODIUM ACETATE. PH

REMARK 280 5.0. VAPOR DIFFUSION. SITTING DROP. TEMPERATURE 294.15K

REMARK 290

REMARK 290 CRYSTALLOGRAPHIC SYMMETRY

REMARK 290 SYMMETRY OPERATORS FOR SPACE GROUP: P 31 2 1

REMARK 290

REMARK 290 SYMOP SYMMETRY

REMARK 290 NNNMM Operator

REMARK 200 1555 X, Y, Z

REMARK 290 2555 -Y, X-Y, Z + 1/3

REMARK 290 3555 -X + Y, -X, Z + 2/3

REMARK 200 4555 Y, X, -Z

REMARK 290 5555 X-Y, -Y, -Z + 2/3

REMARK 290 6555 -X, -X + Y, -Z + 1/3

REMARK 290

REMARK 290 WHERE NMN -> OPERATOR NUMBER

REMARK 290 MMM -> TRANSLATION VECTOR

REMARK 290

REMARK 290 CRYSTALLOGRAPHIC SYMMETRY TRANSFORMATIONS

REMARK 290 THE FOLLOWING TRANSFORMATIONS OPERATE ON THE ATOM/HETATM

REMARK 290 RECORDS IN THIS ENTRY TO PRODUCE CRYSTALLOGRAPHICALLY

REMARK 290 RELATED MOLECULES.

REMARK 290 SMTRY1 1 1.000000 0.000000 0.000000 0.000000

TABLE 3-continued

---

REMARK 290 SMTRY2 1 0.000000 1.000000 0.000000 0.000000

REMARK 290 SMTRY3 1 0.000000 0.000000 1.000000 0.000000

REMARK 290 SMTRY1 2 -0.500000 -0.866025 0.000000 0.000000

REMARK 290 SMTRY2 2 0.866025 -0.500000 0.000000 0.000000

REMARK 290 SMTRY3 2 0.000000 0.000000 1.000000 27.49067

REMARK 290 SMTRY1 3 -0.500000 0.866025 0.000000 0.000000

REMARK 290 SMTRY2 3 -0.866025 -0.500000 0.000000 0.000000

REMARK 290 SMTRY3 3 0.000000 0.000000 1.000000 54.98133

REMARK 290 SMTRY1 4 -0.500000 0.866025 0.000000 0.000000

REMARK 290 SMTRY2 4 0.866025 0.500000 0.000000 0.000000

REMARK 290 SMTRY3 4 0.000000 0.000000 -1.000000 0.000000

REMARK 290 SMTRY1 5 1.000000 0.000000 0.000000 0.000000

REMARK 290 SMTRY2 5 0.000000 -1.000000 0.000000 0.000000

REMARK 290 SMTRY3 5 0.000000 0.000000 -1.000000 54.98133

REMARK 290 SMTRY1 6 -0.500000 -0.866025 0.000000 0.000000

REMARK 290 SMTRY2 6 -0.866025 0.500000 0.000000 0.000000

REMARK 290 SMTRY3 6 0.000000 0.000000 -1.000000 27.49067

REMARK 290

REMARK 290 REMARK: NULL

REMARK 300

REMARK 300 BIOMOLECULE: 1

REMARK 300 SEE REMARK 350 FOR THE AUTHOR PROVIDED AND/OR PROGRAM

REMARK 300 GENERATED ASSEMBLY INFORMATION FOR THE STRUCTURE IN

REMARK 300 THIS ENTRY. THE REMARK MAY ALSO PROVIDE INFORMATION ON

REMARK 300 BURIED SURFACE AREA.

REMARK 350

REMARK 350 COORDINATES FOR A COMPLETE MULTIMER REPRESENTING THE KNOWN

REMARK 350 BIOLOGICALLY SIGNIFICANT OLIGOMERIZATION STATE OF THE

REMARK 350 MOLECULE CAN BE GENERATED BY APPLYING BIOMT TRANSFORMATIONS

REMARK 350 GIVEN BELOW. BOTH NON-CRYSTALLOGRAPHIC AND

REMARK 350 CRYSTALLOGRAPHIC OPERATIONS ARE GIVEN.

REMARK 350

REMARK 350 BIOMOLECULE: 1

REMARK 350 AUTHOR DETERMINED BIOLOGICAL UNIT: DIMERIC

REMARK 350 SOFTWARE DETERMINED QUATERNARY STRUCTURE: DIMERIC

REMARK 350 SOFTWARE USED: PISA

REMARK 350 TOTAL BURIED SURFACE AREA: 4850 ANGSTROM\*\*2

REMARK 350 SURFACE AREA OF THE COMPLEX: 21940 ANGSTROM\*\*2

REMARK 350 CHANGE IN SOLVENT FREE ENERGY: -45.0 KCAL/MOL

REMARK 350 APPLY THE FOLLOWING TO CHAINS: A. B

TABLE 3-continued

REMARK 350 BIOMT1 1 1.000000 0.000000 0.000000 0.000000

REMARK 350 BIOMT2 1 0.000000 1.000000 0.000000 0.000000

REMARK 350 BIOMT3 1 0.000000 0.000000 1.000000 0.000000

REMARK 465

REMARK 465 MISSING: RESIDUES

REMARK 465 THE FOLLOWING RESIDUES WERE NOT LOCATED IN THE

REMARK 465 EXPERIMENT. (M = MODEL NUMBER; RES = RESIDUE NAME; C= CHAIN

REMARK 465 IDENTIFIER, SSSEQ = SEQUENCE NUMBER, I = INSERTION CODE.)

REMARK 465

REMARK 465 M RES C SSSEQI

REMARK 465 MET A 1

REMARK 465 GLY A 2

REMARK 465 SER A 3

REMARK 465 PRO A 4

REMARK 465 LYS A 5

REMARK 465 GLU A 82

REMARK 465 SER A 83

REMARK 465 ARG A 84

REMARK 465 GLN A 85

REMARK 465 GLN A 86

REMARK 465 ASN A 87

REMARK 465 HIS A 88

REMARK 465 ILE A 89

REMARK 465 GLU A 90

REMARK 465 MET B 1

REMARK 465 GLY B 2

REMARK 465 SER B 3

REMARK 465 PRO B 4

REMARK 465 LYS B 5

REMARK 465 THR B 81

REMARK 465 GLU B 82

REMARK 465 SER B 83

REMARK 465 ARG B 84

REMARK 465 GLN B 85

REMARK 465 GLN B 86

REMARK 465 ASN B 87

REMARK 465 HIS B 88

REMARK 465 ILE B 89

REMARK 465 GLU B 90

REMARK 465 GLY B 91

## TABLE 3-continued

REMARK 465 ASN B 278

REMARK 500

REMARK 500 GEOMETRY AND STEREOCHEMISTRY

REMARK 500 SUBTOPIC: CLOSE CONTACTS IN SAME ASYMMETRIC UNIT

REMARK 500

REMARK 500 THE FOLLOWING ATOMS ARE IN CLOSE CONTACT.

REMARK 500

REMARK 500 ATM1 RES C SSEQI ATM2 RES C SSEQI DISTANCE

REMARK 500 O HOH A 1189 O HOH A 1200 2.18

REMARK 500 O HOH B 1186 O HOH B 1205 2.19

REMARK 500

REMARK 500 REMARK: NULL

REMARK 500

REMARK 500 GEOMETRY AND STEREOCHEMISTRY

REMARK 500 SUBTOPIC: COVALENT BOND LENGTHS

REMARK 500

REMARK 500 THE STEREOCHEMICAL PARAMETERS OF THE FOLLOWING RESIDUES

REMARK 500 HAVE VALUES WHICH DEVIATE FROM EXPECTED VALUES BY MORE

REMARK 500 THAN 6\*RMSD (M = MODEL NUMBER: RES = RESIDUE NAME: C = CHAIN

REMARK 500 IDENTIFIER; SSEQ = SEQUENCE NUMBER: I = INSERTION CODE)

REMARK 500

REMARK 500 STANDARD TABLE:

REMARK 500 FORMAT: (10X, I3, 1X.2(A3, 1X, A1, I4, A1, 1X, A4, 3X), 1X, F6.3)

REMARK 500

REMARK 500 EXPECTED VALUES PROTEIN: ENGH AND HUBER. 1999

REMARK 500 EXPECTED VALUES NUCLEIC ACID: CLOWNEY ET AL 1996

REMARK 500

REMARK 500 M RES CSSEQI ATM1 RES CSSEQI ATM2 DEVIATION

REMARK 500 HIS A 34 CG HIS A 34 CD2 0.064

REMARK 500 GLU A 160 C GLU A 160 O 0.118

REMARK 500 HIS A 192 CG HIS A 192 CD2 0.055

REMARK 500 THR B 220 CB THR B 220 CG2 -0.202

REMARK 500

REMARK 500 REMARK: NULL

REMARK 500

REMARK 500 GEOMETRY AND STEREOCHEMISTRY

REMARK 500 SUBTOPIC: COVALENT BOND ANGLES

REMARK 500

REMARK 500 THE STEREOCHEMICAL PARAMETERS OF THE FOLLOWING RESIDUES

REMARK 500 HAVE VALUES WHICH DEVIATE FROM EXPECTED VALUES BY MORE

TABLE 3-continued

REMARK 500 THAN 6\*RMSD (M = MODEL NUMBER: RES = RESIDUE NAME: C = CHAIN

REMARK 500 IDENTIEIER; SSEQ = SEQUENCE NUMBER: I = INSERTION CODE).

REMARK 500

REMARK 500 STANDARD TABLE:

REMARK 500 FORMAT: (10X, I3, 1X, A3, 1X, A1, I4, A1.3

(1X, A4, 2X), I2X, F5.1)

REMARK 500

REMARK 500 EXPECTED VALUES PROTEIN: ENGH AND HUBER. 1999

REMARK 500 EXPECTED VALUES NUCLEIC ACID: CLOWNEY ET AL 1996

REMARK 500

REMARK 500 M RES CSSEQI ATM1 ATM2 ATM3

REMARK 500 ASP A 161 CB - CG - OD1 ANGL. DEV. = 6.7 DEGREES

REMARK 500 ARG A 171 NE - CZ - NH1 ANGL. DEV. = 4.1 DEGREES

REMARK 500 ARG A 171 NE - CZ - NH2 ANGL. DEV. = -3.1 DEGREES

REMARK 500 ARG A 227 NE - CZ - NH1 ANGL. DEV. = 3.5 DEGREES

REMARK 500 ASP A 248 CB - CG - OD2 ANGL. DEV. = -5.8 DEGREES

REMARK 500 ARG S 227 NE - CZ - NH2 ANGL. DEV. = -4.0 DEGREES

REMARK 500

REMARK 500 REMARK: NULL

REMARK 500

REMARK 500 GEOMETRY AND STEREOCHEMISTRY

REMARK 500 SUBTOPIC: TORSION ANGLES

REMARK 500

REMARK 500 TORSION ANGLES OUTSIDE THE EXPECTED RAMACHANDRAN REGIONS:

REMARK 500 (M = MODEL NUMBER; RES = RESIDUE NAME: C = CHAIN IDENTIFIER:

REMARK 500 SSEQ = SEQUENCE NUMBER: I = INSERTION CODE).

REMARK 500

REMARK 500 STANDARD TABLE:

REMARK 500 FORMAT: (10X, I3, 1X, A3, 1X, A1, I4, A1, 4X, F7.2, 3X, F7.2)

REMARK 500

REMARK 500 EXPECTED VALUES: G J KLEYWEGT AND T A JONES 1996). PHI/PSI-

REMARK 500 CHOLOGY. RAMACHANDRAN REVISITED. STRUCTURE 4. 1395-1400

REMARK 500

REMARK 500 M RES CSSEQI PSI PHI

REMARK 500 ARG A 227 81.53 63.17

REMARK 500 PRO A 259 -19.64 -49.20

REMARK 500 THR B 50 -143.74 -116.57

REMARK 500 ASP B 201 31.72 70.03

REMARK 500 ARG B 227 76.48 65.91

REMARK 500

## TABLE 3-continued

REMARK 500 REMARK: NULL

REMARK 500

REMARK 500 GEOMETRY AND STEREOCHEMISTRY

REMARK 500 SUBTOPIC: NON-CIS. NON-TRANS

REMARK 500

REMARK 500 THE FOLLOWING PEPTIDE BONDS DEVIATE SIGNIFICANTLY FROM BOTH

REMARK 500 CIS AND TRANS CONFORMATION. CIS BONDS, IF ANY, ARE LISTED

REMARK 500 ON CISPEP RECORDS. TRANS IS DEFINED AS 180 +/- 30 AND

REMARK 500 CIS IS DEFINED AS 0 +/- 30 DEGREES.

REMARK 500 MODEL OMEGA

REMARK 500 ALA A 277 ASN A 276 -135.53

REMARK 500

REMARK 500 REMARK: NULL

REMARK 500

REMARK 500 GEOMETRY AND STEREOCHEMISTRY

REMARK 500 SUBTOPIC: CHIRAL CENTERS

REMARK 500

REMARK 500 UNEXPECTED CONFIGURATION OF THE FOLLOWING CHIRAL

REMARK 500 CENTER(S) USING IMPROPER CA--C--CB--N CHIRALITY

REMARK 500 FOR AMINO ACIDS AND C1'--O4'--N1(N9)--C2' FOR

REMARK 500 NUCLEIC ACIDS OR EQUIVALENT ANGLE

REMARK 500 M = MODEL NUMBER: RES = RESIDUE NAME: C = CHAIN

REMARK 500 IDENTIFIER: SSEQ = SEQUENCE NUMBER: I = INSERTION CODE

REMARK 500

REMARK 500 STANDARD TABLE:

REMARK 500 FORMAT: (10X, I3, 1X, A3, 1X, A1, I4, A1, 6X, F5.1,

6X, A1, 10X, A1, 3X, A16)

REMARK 500

REMARK 500 M RES CSSEQI IMPROPER EXPECTED FOUND DETAILS

REMARK 500 VAL B 25 24.5 L L OUTSIDE RANGE

REMARK 500 SER B 216 24.9 L L OUTSIDE RANGE

REMARK 500

REMARK 500 REMARK: NULL

REMARK 800

REMARK 800 SITE

REMARK 800 SITE\_IDENTIFIER: AC1

REMARK 800 EVIDENCE\_CODE: SOFTWARE

REMARK 800 SITE\_DESCRIPTION: BINDING SITE FOR RESIDUE BMX A 1000

REMARK 800

REMARK 800 SITE\_IDENTIFIER: AC2

TABLE 3-continued

REMARK 800 EVIDENCE\_CODE: SOFTWARE

REMARK 800 SITE\_DESCRIPTION: BINDING SITE FOR RESIDUE BMX B 1000

DBREF 4IVN A I 278 UNP Q7MD38 Q7MD38\_VIBVY 1 278

DBREF 4IVN B 1 278 UNP Q7MD38 Q7MD38\_VIBVY 1 278

SEQRES 1 A 278 MET GLY SER PRO LYS ASN LEU LEU VAL ARG LEU ARG SER

SEQRES 2 A 278 ASN MET GLU PRO PHE SER LYS LYS LEU ARG VAL VAL ALA

SEQRES 3 A 278 ASP TYR ILE LEU GLU ASN ALA HIS ASP VAL GLN PHE GLN

SEQRES 4 A 278 THR ILE THR ASP LEU ALA ARG ASN THR GLN THR SER GLU

SEQRES 5 A 278 ALA THR VAL VAL ARG LEU CYS ARG ASP MET GLY TYR LYS

SEQRES 6 A 278 GLY TYR SER ASP PHE ARG MET ALA LEU ALA VAL ASP LEU

SEQRES 7 A 278 SER GLN THR GLU SER ARG GLN ASN HIS ILE GLU GLY

SEQRES 8 A 278 ASP ILE CYS ASP VAL SER ALA GLN SER ALA VAL ASP SER

SEQRES 9 A 278 LEU GLN ASP THR ALA LYS LEU ILE ASP ARG LYS SER LEU

SEQRES 10 A 278 ALA ARG ILE VAL GLU ARG VAL HIS GLN ALA GLU PHE ILE

SEQRES 11 A 278 GLY CYS ILE GLY VAL GLY ALA SER SER ILE VAL GLY ARG

SEQRES 12 A 278 TYR LEU ALA TYR ARG LEU ILE ARG ILE GLY LYS ALA

SEQRES 13 A 278 ILE MET PHE GLU ASP THR HIS LEU ALA ALA MET SER ALA

SEQRES 14 A 278 SER ARG SER SER GLN GLY ASP LEU TRP PHE ALA VAL SER

SEQRES 15 A 278 SER SER GLY SER THR LYS GLU VAL ILE HIS ALA ALA GLY

SEQRES 16 A 278 LEU ALA TYR LYS ARG ASP ILE PRO VAL VAL SER LEU THR

SEQRES 17 A 278 ASN ILE ASN HIS SER PRO LEU SER SER LEU SEE THR GLU

SEQRES 18 A 278 MET LEU VAL ALA ALA ARG PRO GLU GLY PRO LEU THR GLY

SEQRES 19 A 278 GLY ALA PHE ALA SER LYS VAL GLY ALA LEU LEU LEU VAL

SEQRES 20 A 278 ASP VAL LEU VAL ASN SER LEU LEU GLU SER TYR PRO GLU

SEQRES 21 A 278 TYR LYS ASP SER VAL GLN GLU THR ALA GLU VAL VAL ILE

SEQRES 22 A 278 PRO LEU MET ALA ASN

SEQRES 1 B 278 MET GLY SER PRO LYS ASN LEU LEU VAL ARG LEU ARG SER

SEQRES 2 B 278 ASN MET GLU PRO PHE SER LYS LYS LEU ARG VAL VAL ALA

SEQRES 3 B 278 ASP TYR ILE LEU GLU ASN ALA HIS ASP VAL GLN PHE GLN

SEQRES 4 B 278 THR ILE THR ASP LEU ALA ARG ASN THR GLN THR SER GLU

SEQRES 5 B 278 ALA THR VAL VAL ARG LEU CYS ARG ASP MET GLY TYR LYS

SEQRES 6 B 278 GLY TYR SER ASP PHE ARG MET ALA LEU ALA VAL ASP LEU

SEQRES 7 B 278 SER GLN THR GLU SER ARG GLN GLN ASN HIS ILE GLU GLY

SEQRES 8 B 278 ASP ILE CYS ASP VAL SER ALA GLN SER ALA VAL ASP SER

SEQRES 9 B 278 LEU GLN ASP THR ALA LYS LEU ILE ASP ARG LYS SER LEU

SEQRES 10 B 278 ALA ARG ILE VAL GLU ARG VAL HIS GLN ALA GLU PHE ILE

SEQRES 11 B 278 GLY CYS ILE GLY VAL GLY ALA SER SER ILE VAL GLY ARG

SEQRES 12 B 278 TYR LEU ALA TYR ARG LEU ILE ARG ILE GLY LYS LYS ALA

SEQRES 13 B 278 ILE MET PHE GLU ASP THR HIS LEU ALA ALA MET SER ALA

SEQRES 14 B 278 SER ARG SER SER GLN GLY ASP LEU TRP PHE ALA VAL SER

TABLE 3-continued

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SEQRES 15 B 278 SER SER GLY SER THR LYS GLU VAL ILE HIS ALA ALA GLY  
 SEQRES 16 B 278 LEU ALA TYR LYS ARG ASP ILE PRO VAL VAL SER LEU THR  
 SEQRES 17 B 278 ASN ILE ASN HIS SER PRO LEU SER SER LEU SER THR GLU  
 SEQRES 18 B 278 MET LEU VAL ALA ALA ARG PRO GLU GLY PRO LEU THR GLY  
 SEQRES 19 B 278 GLY ALA PHE ALA SER LYS VAL GLY ALA LEU LEU LEU VAL  
 SEQRES 20 B 278 ASP VAL LEU VAL ASN SER LEU LEU GLU SER TYR PRO GLU  
 SEQRES 21 B 278 TYR LYS ASP SER VAL GLN GLU THR ALA GLU VAL VAL ILE  
 SEQRES 22 B 278 PRO LEU MET ALA ASN  
  
 HET BMX A1000 19  
 HET BMX B1000 19  
  
 HETNAM BMX 2-(ACETYLAMINO)-2-DEOXY-5-O-PHOSPHONO-ALPHA-D-  
 HETNAM 2 BMX MANNOPYRANOSE  
  
 FORMUL 3 BMX 2(CB H16 N 09 P)  
 FORMUL 5 HOH \*221(H2 O)  
  
 HELIX 1 1 ASN A 6 ASN A 14 1 9  
 HELIX 2 2 SER A 19 GLN A 37 1 19  
 HELIX 3 3 THR A 40 GLN A 49 1 10  
 HELIX 4 4 SER A 51 MET A 62 1 12  
 HELIX 5 5 GLY A 66 SER A 79 1 14  
 HELIX 6 6 ASP A 92 ILE A 112 1 21  
 HELIX 7 7 ASP A 113 ALA A 127 1 15  
 HELIX 8 8 GLY A 136 ILE A 152 1 17  
 HELIX 9 9 ASP A 161 ARG A 171 1 11  
 HELIX 10 10 THR A 187 ARG A 200 1 14  
 HELIX 11 11 LEU A 215 SER A 219 5 5  
 HELIX 12 12 ALA A 236 TYR A 258 1 23  
 HELIX 13 13 GLU A 260 VAL A 271 1 12  
 HELIX 14 14 VAL A 272 MET A 276 5 5  
 HELIX 15 15 LEU B 7 ASN B 14 1 8  
 HELIX 16 16 SER B 19 GLN B 37 1 19  
 HELIX 17 17 THR B 40 GLN B 49 1 10  
 HELIX 18 18 SER B 51 MET B 62 1 12  
 HELIX 19 19 GLY B 66 GLN B 80 1 15  
 HELIX 20 20 ILE B 93 ILE B 112 1 20  
 HELIX 21 21 ASP B 113 ALA B 127 1 15  
 HELIX 22 22 GLY B 136 ILE B 152 1 17  
 HELIX 23 23 ASP B 161 ARG B 171 1 11  
 HELIX 24 24 THR B 187 ARG B 200 1 14  
 HELIX 25 25 SER B 213 SER B 219 5 7  
 HELIX 26 26 ALA B 236 TYR B 258 1 23

TABLE 3-continued

HELIX 27 27 GLU B 260 VAL B 271 1 12  
 HELIX 28 28 VAL B 272 MET B 276 5 5  
 SHEET 1 A 5 ALA A 156 PHE A 159 0  
 SHEET 2 A 5 PHE A 129 ILE A 133 1 N CYS A 132 O ILE A 157  
 SHEET 3 A 5 ASP A 176 VAL A 181 1 O VAL A 181 N ILE A 133  
 SHEET 4 A 5 VAL A 204 THR A 208 1 O VAL A 205 N TRP A 178  
 SHEET 5 A 5 GLU A 221 VAL A 224 1 O LEU A 223 N SER A 206  
 SHEET 1 B 5 LYS B 155 PHE B 169 0  
 SHEET 2 B 5 PHE B 129 ILE B 133 1 N CYS B 132 O PHE B 150  
 SHEET 3 B 5 ASP B 176 VAL B 181 1 O VAL B 181 N ILE B 133  
 SHEET 4 B 5 VAL B 204 THR B 208 1 O VAL B 205 N TRP B 178  
 SHEET 5 B 5 GLU B 221 VAL B 224 1 O LEU B 223 N SER B 206  
 SITE 1 AC1 19 VAL A 135 ALA A 137 SER A 138 HIS A 163  
 SITE 2 AC1 19 MET A 167 SER A 182 SER A 183 SER A 184  
 SITE 3 AC1 19 THR A 187 GLY A 230 HOH A1102 HOH A1103  
 SITE 4 AC1 19 HOH A1105 HOH A1108 HOH A1109 TYR B 147  
 SITE 5 AC1 19 ARG B 151 VAL B 272 MET B 276  
 SITE 1 AC2 19 TYR A 147 ARG A 151 VAL A 272 VAL B 135  
 SITE 2 AC2 19 ALA B 137 SER B 138 HIS B 163 MET B 167  
 SITE 3 AC2 19 SER B 182 SER B 183 SER B 184 THR B 187  
 SITE 4 AC2 19 GLY B 230 GLY B 234 HOH B1102 HOH B1104  
 SITE 5 AC2 19 HOH B1113 HOH B1116 HOH B1119  
 CRYST1 109.205 109.205 82.472 90.00 90.00 120.00 P 31 2 1 12  
 ORIGX1 1.000000 0.000000 0.000000 0.00000  
 ORIGX2 0.000000 1.000000 0.000000 0.00000  
 ORIGX3 0.000000 0.000000 1.000000 0.00000  
 SCALE1 0.009157 0.005287 0.000000 0.00000  
 SCALE2 0.000000 0.010574 0.000000 0.00000  
 SCALE3 0.000000 0.000000 0.012125 0.00000  
 ATOM 1 N ASN A 6 18.407 -18.282 7.841 1.00 65.46 N  
 ANISOU 1 N ASN A 6 8081 7427 9362 -71 -1349 -1116 N  
 ATOM 2 CA ASN A 6 17.833 -18.480 9 220 1.00 78.17 C  
 ANISOU 2 CA ASN A 6 9687 8914 11100 -298 -1254 -922 C  
 ATOM 3 C ASN A 6 17.340 -17.238 9.992 1.00 70.53 C  
 ANISOU 3 C ASN A 6 8645 8122 10032 -404 -1084 -781 C  
 ATOM 4 O ASN A 6 16.118 -17.096 10.194 1.00 66.38 O  
 ANISOU 4 O ASN A 6 8016 7595 9612 -550 -1052 -786 O  
 ATOM 5 CB ASN A 6 18.736 -19.329 10.109 1.00 88.32 C  
 ANISOU 5 CB ASN A 6 11096 10027 12435 -313 -1258 -782 C  
 ATOM 6 CG ASN A 6 17.953 -20.005 11.207 1.00 95.86 C

TABLE 3-continued

ANISOU 6 CG ASN A 6	12184 10920 13697 -546 -1219 -623 C
ATOM 7 OD1 ASN A 6	17.873 -21.226 11.251 1.00 107.17 O
ANISOU 7 OD1 ASN A 6	13547 11970 15202 -597 -1328 -633 O
ATOM 8 ND2 ASN A 6	17.322 -19.210 12.071 1.00 97.16 N
ANISOU 8 ND2 ASN A 6	12156 11085 13676 -693 -1062 -481 N
ATOM 9 N LEU A 7	18.256 -16.363 10.444 1.00 58.71 N
ANISOU 9 N LEU A 7	7186 5771 8352 -334 -981 -670 N
ATOM 10 CA LEU A 7	17.874 -14.959 10.576 1.00 53.24 C
ANISOU 10 CA LEU A 7	6409 6277 7542 -347 -868 -634 C
ATOM 11 C LEU A 7	17.247 -14.501 9.231 1.00 52.37 C
ANISOU 11 C LEU A 7	6217 6266 7416 -252 -947 -805 C
ATOM 12 O LEU A 7	16.236 -13.769 9.229 1.00 51.26 O
ANISOU 12 O LEU A 7	5972 6208 7298 -314 -908 -823 O
ATOM 13 CB LEU A 7	19.043 -14.019 10.921 1.00 50.90 C
ANISOU 13 CB LEU A 7	6155 6120 7063 -254 -782 -540 C
ATOM 14 CG LEU A 7	18.713 -12.520 10.859 1.00 49.25 C
ANISOU 14 CG LEU A 7	5865 6091 6757 -243 -693 -528 C
ATOM 15 CD1 LEU A 7	19.803 -11.693 11.514 1.00 47.98 C
ANISOU 15 CD1 LEU A 7	5741 6023 6467 -204 -605 -420 C
ATOM 16 CD2 LEU A 7	18.585 -12.057 9.419 1.00 55.09 C
ANISOU 16 CD2 LEU A 7	6565 6925 7442 -106 -766 -653 C
ATOM 17 N LEU A 8	17.826 -14.909 8.095 1.00 48.71 N
ANISOU 17 N LEU A 8	5796 5808 6902 -89 -1061 -936 N
ATOM 18 CA LEU A 8	17.236 -14.480 6.813 1.00 52.64 C
ANISOU 18 CA LEU A 8	6227 6424 7349 14 -1145 -1090 C
ATOM 19 C LEU A 8	15.872 -15.125 6.516 1.00 54.82 C
ANISOU 19 C LEU A 8	6418 6600 7813 -84 -1252 -1224 C
ATOM 20 O LEU A 8	14.938 -14.452 5.025 1.00 53.15 O
ANISOU 20 O LEU A 8	6104 6493 7596 -80 -1276 -1290 O
ATOM 21 CB LEU A 8	18.198 -14.636 5.632 1.00 56.78 C
ANISOU 21 CB LEU A 8	6811 7036 7725 226 -1224 -1200 C
ATOM 22 CG LEU A 8	19.337 -13.604 5.585 1.00 59.35 C
ANISOU 22 CG LEU A 8	7168 7535 7847 331 -1112 -1082 C
ATOM 23 CD1 LEU A 8	20.359 -14.019 4.548 1.00 61.43 C
ANISOU 23 CD1 LEU A 8	7482 7879 7980 527 -1172 -1189 C
ATOM 24 CD2 LEU A 8	18.807 -12.203 5.309 1.00 61.52 C
ANISOU 24 CD2 LEU A 8	7375 7970 8028 336 -1051 -1023 C
ATOM 25 N VAL A 9	15.746 -15.410 6.841 1.00 55.47 N
ANISOU 25 N VAL A 9	6532 6467 8078 -175 -1322 -1259 N
ATOM 26 CA VAL A 9	14.429 -17.074 6.760 1.00 58.23 C

TABLE 3-continued

ANISOU 26 CA VAL A 9	6780 6689 8654 -317 -1409 -1365 C
ATOM 27 C VAL A 9	13.437 -16.415 7.731 1.00 57.99 C
ANISOU 27 C VAL A 9	6635 6706 8693 -504 -1271 -1237 C
ATOM 28 O VAL A 9	12.280 16.188 7.358 1.00 57.55 O
ANISOU 28 O VAL A 9	6441 5695 8730 -557 -1316 -1342 O
ATOM 29 CB VAL A 9	14.474 -18.614 7.032 1.00 57.89 C
ANISOU 29 CB VAL A 9	6796 6362 8839 -410 -1507 -1399 C
ATOM 30 CG1 VAL A 9	13.036 -19.236 6.987 1.00 59.10 C
ANISOU 30 CG1 VAL A 9	5817 6379 9260 -593 -1588 -1499 C
ATOM 31 CG2 VAL A 9	15.470 -19.286 6.080 1.00 54.95 C
ANISOU 31 CG2 VAL A 9	6529 5945 8405 -202 -1649 -1558 C
ATOM 32 N ARG A 10	13.879 -16.101 8.950 1.00 60.02 N
ANISOU 32 N ARG A 10	6938 5969 8896 -592 -1109 -1030 N
ATOM 33 CA ARG A 10	12.962 -15.536 9.943 1.00 59.43 C
ANISOU 33 CA ARG A 10	6753 6954 8872 -762 -966 -924 C
ATOM 34 C ARG A 10	12.503 -14.194 9.486 1.00 63.36 C
ANISOU 34 C ARG A 10	7151 7660 9262 -673 -937 -984 C
ATOM 35 O ARG A 10	11.310 -13.887 9.640 1.00 63.30 O
ANISOU 35 O ARG A 10	6991 7700 9361 -772 -910 -1026 O
ATOM 36 CB ARG A 10	13.483 -15.574 11.378 1.00 60.57 C
ANISOU 36 CB ARG A 10	6973 7073 8969 -872 -810 -704 C
ATOM 37 CG ARG A 10	13.546 -17.002 11.931 1.00 65.64 C
ANISOU 37 CG ARG A 10	7688 7478 9776 -1007 -843 -620 C
ATOM 38 CD ARG A 10	13.686 -17.062 13.447 1.00 73.54 C
ANISOU 38 CD ARG A 10	8733 8471 10736 -1155 -681 -384 C
ATOM 39 NE ARC A 10	14.758 -16.175 13.887 1.00 83.65 N
ANISOU 39 NE ARG A 10	10100 9901 11784 -1032 -604 -301 N
ATOM 40 CZ ARG A 10	14.600 -14.871 14.134 1.00 85.86 C
ANISOU 40 CZ ARG A 10	10308 10389 11926 -998 -497 -305 C
ATOM 41 NH1 ARG A 10	13.408 -14.290 14.008 1.00 89.12 N
ANISOU 41 NH1 ARG A 10	10562 10898 12401 -1065 -449 -383 N
ATOM 42 NH2 ARG A 10	15.640 -14.143 14.496 1.00 80.60 N
ANISOU 42 HH2 ARG A 10	9719 9828 11079 -893 -449 -243 N
ATOM 43 N LEU A 11	13.396 -13.442 8.823 1.00 61.15 N
ANISOU 43 N LEU A 11	6945 7497 8793 -484 -958 -996 N
ATOM 44 CA LEU A 11	13.060 -12.083 8.336 1.00 57.08 C
ANISOU 44 CA LEU A 11	6356 7160 8172 -383 -941 -1026 C
ATOM 45 C LEU A 11	11.935 -12.184 7.310 1.00 62.42 C
ANISOU 45 C LEU A 11	6914 7862 8941 -350 -1083 -1203 C
ATOM 46 O LEU A 11	10.889 -11.519 7.422 1.00 64.23 O

TABLE 3-continued

ANISOU 46 O LEU A 11	7004 8166 9235 -392 -1064 -1235 O
ATOM 47 CB LEU A 11	14.281 -11.356 7.750 1.00 51.02 C
ANISOU 47 CB LEU A 11	5694 6496 7197 -202 -938 -983 C
ATOM 48 CG LEU A 11	14.332 -9.827 7.472 1.00 48.59 C
ANISOU 48 CG LEU A 11	5355 6343 6764 -103 -890 -936 C
ATOM 49 CD1 LEU A 11	13.734 -8.925 8.551 1.00 47.87 C
ANISOU 49 CD1 LEU A 11	5180 6286 6723 -204 -766 -554 C
ATOM 50 CD2 LEU A 11	15.773 -9.397 7.195 1.00 47.01 C
ANISOU 50 CD2 LEU A 11	5268 6201 6391 18 -854 -849 C
ATOM 51 N ARG A 12	12.177 -13.000 6.304 1.00 62.22 N
ANISOU 51 N ARG A 12	6940 7784 8916 -260 -1236 -1333 N
ATOM 52 CA ARG A 12	11.161 -13.339 5.313 1.00 65.79 C
ANISOU 52 CA ARG A 12	7414 8371 9592 -229 -1406 -1532 C
ATOM 53 C ARG A 12	9.747 -13.675 5.870 1.00 67.06 C
ANISOU 53 C ARG A 12	7271 8334 9875 -427 -1403 -1582 C
ATOM 54 O ARG A 12	8.751 -13.026 5.515 1.00 61.31 O
ANISOU 54 O ARG A 12	6398 7712 9184 -407 -1448 -1664 O
ATOM 55 CB ARG A 12	11.668 -14.512 4.477 1.00 58.50 C
ANISOU 55 CB ARG A 12	7719 8488 9821 -148 -1562 -1678 C
ATOM 56 CG ARG A 12	12.419 14.097 3.240 1.00 75.79 C
ANISOU 56 CG ARG A 12	8726 9567 10502 94 -1645 -1751 C
ATOM 57 CD ARG A 12	11.467 -13.429 2.259 1.00 83.06 C
ANISOU 57 CD ARG A 12	9543 10643 11373 192 -1765 -1875 C
ATOM 58 NE ARG A 12	10.071 -13.817 2.471 1.00 86.82 N
ANISOU 58 NE ARG A 12	9855 11037 12095 45 -1841 -1987 N
ATOM 59 CZ ARG A 12	9.063 -13.495 1.655 1.00 94.85 C
ANISOU 59 CZ ARG A 12	10751 12158 13130 111 -1987 -2137 C
ATOM 60 NH1 ARG A 12	9.288 -12.784 0.554 1.00 95.02 N
ANISOU 60 NH1 ARG A 12	10819 12358 12918 329 -2075 -2177 N
ATOM 61 NH2 ARG A 12	7.824 -13.893 1.931 1.00 93.22 N
ANISOU 61 NH2 ARG A 12	10371 11873 13174 -42 -2046 -2241 N
ATOM 62 N SER A 13	9.670 -14.704 6.712 1.00 67.77 N
ANISOU 62 N SER A 13	7366 8244 10138 -613 -1355 -1528 N
ATOM 63 CA SER A 13	8.381 -15.232 7.120 1.00 76.61 C
ANISOU 63 CA SER A 13	8312 9283 11514 -816 -1359 -1580 C
ATOM 64 C SER A 13	7.775 -14.367 8.210 1.00 76.77 C
ANISOU 64 C SER A 13	8216 9413 11542 -935 -1164 -1448 C
ATOM 65 O SER A 13	6.606 -14.025 8.129 1.00 79.19 O
ANISOU 65 O SER A 13	8326 9795 11967 -990 -1176 -1538 O
ATOM 66 CB SER A 13	8.434 -16.730 7.498 1.00 78.82 C

TABLE 3-continued

ANISOU 66 CB SER A 13	8635 9307 12006 -982 -1399 -1569 C
ATOM 67 CG SER A 13	9.521 -17.045 8.351 1.00 73.03 O
ANISOU 67 CG SER A 13	8072 8485 11192 -1009 -1285 -1375 O
ATOM 68 N ASN A 14	8.568 -13.947 9.191 1.00 76.90 N
ANISOU 68 N ASN A 14	8338 9458 11421 -952 -992 -1257 N
ATOM 69 CA ASN A 14	7.991 -13.183 10.305 1.00 84.08 C
ANISOU 69 CA ASN A 14	9136 10480 12330 -1063 -802 -1153 C
ATOM 70 C ASN A 14	7.750 -11.709 9.956 1.00 83.18 C
ANISOU 70 C ASN A 14	8950 10557 12097 -907 -792 -1207 C
ATOM 71 O ASDN A 14	7.525 -10.880 10.861 1.00 86.57 O
ANISOU 71 O ASN A 14	9318 11092 12482 -945 -637 -1134 O
ATOM 72 CB ASN A 14	8.801 -13.297 11.628 1.00 85.20 C
ANISOU 72 CB ASN A 14	9408 10595 12376 -1154 -624 -938 C
ATOM 73 CG ASN A 14	9.451 -14.662 11.830 1.00 91.38 C
ANISOU 73 CG ASN A 14	10325 11168 13226 -1237 -666 -856 C
ATOM 74 OD1 ASN A 14	9.088 -15.673 11.203 1.00 91.91 O
ANISOU 74 OD1 ASN A 14	10369 11081 13471 -1287 -803 -955 O
ATOM 75 ND2 ASN A 14	10.448 -14.689 12.714 1.00 97.54 N
ANISOU 75 ND2 ASN A 14	11268 11947 13885 -1242 -664 -681 N
ATOM 76 N MET A 15	7.772 -11.377 8.666 1.00 74.06 N
ANISOU 76 N MET A 15	7805 9447 10889 -728 -961 -1336 N
ATOM 77 CA MET A 15	7.565 -9.982 8.288 1.00 77.63 C
ANISOU 77 CA MET A 15	8204 10052 11238 -574 -968 -1362 C
ATOM 78 C MET A 15	6.144 -9.578 7.896 1.00 78.15 C
ANISOU 78 C MET A 15	8047 10200 11445 -568 -1045 -1509 C
ATOM 79 O MET A 15	5.857 -8.381 7.712 1.00 79.92 O
ANISOU 79 O MET A 15	8214 10538 11613 -444 -1049 -1523 O
ATOM 80 CB MET A 15	8.625 -9.488 7.287 1.00 80.06 C
ANISOU 80 CB MET A 15	8676 10402 11342 -360 -1060 -1346 C
ATOM 81 CG MET A 15	8.410 -9.785 5.813 1.00 86.13 C
ANISOU 81 CG MET A 15	9445 11196 12086 -220 -1273 -1497 C
ATOM 82 SD MET A 15	9.633 -8.823 4.869 1.00 92.69 S
ANISOU 82 SD MET A 15	10448 12136 12634 22 -1806 -1413 S
ATOM 83 CE MET A 15	8.742 -7.282 4.601 1.00 82.39 C
ANISOU 83 CE MET A 15	9022 10958 11324 128 -1333 -1413 C
ATOM 84 N GLU A 16	5.258 -10.571 7.781 1.00 81.86 N
ANISOU 84 N GLU A 16	8364 10602 12117 -703 -1115 -1620 N
ATOM 85 CA GLU A 16	3.818 -10.315 7.647 1.00 77.99 C
ANISOU 85 CA GLU A 16	7637 10190 11804 -742 -1165 -1761 C
ATOM 86 C GLU A 16	3.317 -9.343 8.732 1.00 77.39 C

TABLE 3-continued

ANISOU 86 C GLU A 16	7433 10229 11742 -791 -971 -1696 C
ATOM 87 O GLU A 16	2.746 -8.286 8.401 1.00 69.65 O
ANISOU 87 O GLU A 16	6342 9367 10756 -654 -1019 -1773 O
ATOM 88 CB GLU A 16	3.022 -11.633 7.657 1.00 85.67 C
ANISOU 88 CB GLU A 16	8475 11048 13029 -942 -1224 -1859 C
ATOM 89 CG GLU A 16	3.185 -12.455 6.390 1.00 92.96 C
ANISOU 89 CG GLU A 16	9463 11881 13975 -858 -1469 -2012 C
ATOM 90 CD GLU A 16	2.443 -11.857 5.197 1.00 105.10 C
ANISOU 90 CD GLU A 16	10880 13548 15504 -676 -1680 -2205 C
ATOM 91 OE1 GLU A 16	2.076 -12.639 4.279 1.00 107.96 O
ANISOU 91 OE1 GLU A 16	11197 13856 15965 -659 -1892 -2389 O
ATOM 92 OE2 GLU A 16	2.215 -10.614 5.172 1.00 98.81 O
AISOU 92 OE2 GLU A 16	10034 12901 14609 -543 -1650 -2179 O
ATOM 93 N PRO A 17	3.566 -9.674 10.029 1.00 79.96 N
ANISOU 93 N PRO A 17	7784 10526 12073 -968 -759 -1556 N
ATOM 94 CA PRO A 17	3.139 -8.757 11.101 1.00 82.09 C
ANISOU 94 CA PRO A 17	7936 10928 12326 -1000 -568 -1515 C
ATOM 95 C PRO A 17	3.819 -7.368 11.136 1.00 83.55 C
ANISOU 95 C RRO A 17	8230 11192 12325 -803 -542 -1474 C
ATOM 96 O PRO A 17	3.335 -6.502 11.868 1.00 86.28 O
ANISOU 96 O PRO A 17	8455 11648 12678 -793 -422 -1495 O
ATOM 97 CB PRO A 17	3.469 -9.533 12.395 1.00 83.52 C
ANISOU 97 CB PRO A 17	8173 11063 12499 -1217 -364 -1353 C
ATOM 98 CG PRO A 17	4.501 -10.541 12.014 1.00 83.68 C
ANISOU 98 CG PRO A 17	8415 10909 12470 -1233 -451 -1269 C
ATOM 99 CD PRO A 17	4.198 -10.900 10.582 1.00 80.28 C
ANISOU 99 CD PRO A 17	7954 10417 12131 -1136 -693 -1433 C
ATOM 100 N PHE A 18	4.907 -7.148 10.383 1.00 78.01 N
ANISOU 100 N PHE A 18	7737 10434 11469 -652 -546 -1422 N
ATOM 101 CA PHE A 18	5.588 -5.834 10.389 1.00 71.25 C
ANISOU 101 CA PHE A 18	6981 9628 10464 -485 -625 -1366 C
ATOM 102 C PHE A 18	4.782 -4.762 9.667 1.00 69.19 C
ANISOU 102 C PHE A 18	6590 9442 10257 -319 -741 -1479 C
ATOM 103 O PHE A 18	4.179 -5.039 8.606 1.00 61.10 O
ANISOU 103 O PHE A 18	5495 8423 9299 -253 -917 -1583 O
ATOM 104 CB PHE A 18	6.973 -5.893 9.715 1.00 73.47 C
ANISOU 104 CB PHE A 18	7501 9841 10575 -376 -697 -1268 C
ATOM 105 CG PHE A 18	7.977 -6.778 10.413 1.00 63.72 C
ANISOU 105 CG PHE A 18	8950 11060 11801 -495 -598 -1147 C
ATOM 106 CD1 PHE A 18	7.769 -7.243 11.725 1.00 89.89 C

TABLE 3-continued

ANISOU	106	CD1	PHE	A	18	9687	11840	12627	-674	-434	-1088	C
ATOM	107	CD2	PHE	A	18	9.172	-7.125	9.765	1.00	84.61	C	
ANISOU	107	CD2	PHE	A	18	9249	1111	11787	-413	-669	-1084	C
ATOM	108	CE1	PHE	A	18	8.711	-8.052	12.360	1.00	90.93	C	
ANISOU	108	CE1	PHE	A	18	9969	11894	12687	-765	-363	-961	C
ATOM	109	CE2	PHE	A	18	10.119	-7.936	10.397	1.00	86.75	C	
ANISOU	109	CE2	PHE	A	18	9653	11305	12003	-499	-597	-980	C
ATOM	110	CZ	PHE	A	18	9.891	-8.390	11.701	1.00	94.92	C	
ANISOU	110	CZ	PHE	A	18	10655	12323	13089	-672	-453	-912	C
ATOM	111	N	SER	A	19	4.844	-3.544	10.227	1.00	62.85	N	
ANISOU	111	N	SER	A	19	5771	8688	9420	-240	-660	-1460	N
ATOM	112	CA	SER	A	19	4.353	-2.300	9.614	1.00	66.57	C	
ANISOU	112	CA	SER	A	19	6171	9198	9926	-48	-770	-1527	C
ATOM	113	C	SER	A	19	5.045	-1.954	8.290	1.00	68.58	C	
ANISOU	113	C	SER	A	19	6589	9408	10061	126	-944	-1461	C
ATOM	114	O	SER	A	19	6.112	-2.481	8.003	1.00	63.85	O	
ANISOU	114	O	SER	A	19	6171	8758	9330	108	-942	-1358	O
ATOM	115	CB	SER	A	19	4.565	-1.120	10.575	1.00	66.24	C	
ANISOU	115	CB	SER	A	19	6126	9175	9865	-5	-642	-1506	C
ATOM	116	OG	SER	A	19	5.957	-0.769	10.649	1.00	62.18	O	
ANISOU	116	OG	SER	A	19	5836	8589	9199	29	-608	-1361	O
ATOM	117	N	LYS	A	20	4.447	-1.053	7.503	1.00	69.33	N	
ANISOU	117	N	LYS	A	20	6616	9531	10195	300	-1089	-1513	N
ATOM	118	CA	LYS	A	20	5.043	-0.594	6.238	1.00	68.35	C	
ANISOU	118	CA	LYS	A	20	6645	9388	9937	476	-1246	-1425	C
ATOM	119	C	LYS	A	20	6.471	-0.065	6.440	1.00	66.56	C	
ANISOU	119	C	LYS	A	20	6643	9108	9576	492	-1152	-1244	C
ATOM	120	O	LYS	A	20	7.372	-0.387	5.658	1.00	67.62	O	
ANISOU	120	O	LYS	A	20	6925	9223	9544	534	-1200	-1146	O
ATOM	121	CB	LYS	A	20	4.195	0.478	5	509	1.00	68.61	C
ANISOU	121	CB	LYS	A	20	6582	9450	10036	672	-1410	-1474	C
ATOM	122	CG	LY3	A	20	4.834	0.882	4.174	1.00	69.04	C	
ANISOU	122	CG	LYS	A	20	6810	9502	9919	845	-1564	-1347	C
ATOM	123	CD	LYS	A	20	3.876	1.370	3.103	1.00	72.11	C	
ANISOU	123	CD	LYS	A	20	7111	9949	10337	1035	-1791	-1412	C
ATOM	124	CE	LYS	A	20	4.145	2.828	2.768	1.00	74.09	C	
ANISOU	124	CE	LYS	A	20	7450	10140	10560	1208	-1846	-1263	C
ATOM	125	NZ	LYS	A	20	3.385	3.265	1.568	1.00	77.24	N	
ANISOU	125	NZ	LYS	A	20	7812	10599	10938	1417	-2090	-1280	N
ATOM	126	N	LYS	A	21	6.665	0.764	7.462	1.00	62.17	N	

TABLE 3-continued

ANISOU 126 N LYS A 21	6062 8502 9057 466 -1024 -127 N
ATOM 127 CA LYS A 21	7.940 1.408 7.636 1.00 59.46 C
ANISOU 127 CA LYS A 21	5891 8090 8610 486 -957 -1064 C
ATOM 128 C LYS A 21	9.017 0.317 7.890 1.00 54.84 C
ANISOU 128 C LYS A 21	5437 7500 7901 358 -864 -989 C
ATOM 129 O LYS A 21	10.090 0.292 7.235 1.00 55.38 O
ANISOU 129 O LYS A 21	5658 7549 7833 404 -887 -865 O
ATOM 130 CB LYS A 21	7.864 2.446 8.763 1.00 65.36 C
ANISOU 130 CB LYS A 21	6582 8797 9455 479 -852 -1094 C
ATOM 131 CG LYS A 21	8.880 3.589 8.643 1.00 73.84 C
ANISOU 131 CG LYS A 21	7795 9771 10491 558 -854 -956 C
ATOM 132 CD LYS A 21	8.363 4.894 9.278 1.00 79.58 C
ANISOU 132 CD LYS A 21	8430 10434 11371 635 -850 -1034 C
ATOM 133 CE LLS A 21	9.460 5.752 9.945 1.00 84.28 C
ANISOU 133 CE LYS A 21	9132 10925 11965 613 -772 -957 C
ATOM 134 NZ LYS A 21	10.730 6.111 9.226 1.00 80.19 N
ANISOU 134 NZ LYS A 21	8790 10320 11357 634 -802 -753 N
ATOM 135 N LEU A 22	8.725 -0.583 8.825 1.00 52.25 N
ANISOU 135 N LEU A 22	5041 7193 7619 204 -760 -1057 N
ATOM 136 CA LEU A 22	9.595 -1.736 9.117 1.00 54.36 C
ANISOU 136 CA LEU A 22	5419 7439 7797 87 -692 -996 C
ATOM 137 C LEU A 22	9.814 -2.643 7.867 1.00 53.25 C
ANISOU 137 C LEU A 22	5349 7300 7565 135 -823 -998 C
ATOM 138 O LEU A 22	10.910 -3.151 7.663 1.00 47.30 O
ANISOU 138 O LEU A 22	4734 6522 6716 130 -807 -917 O
ATOM 139 CB LEU A 22	9.077 -2.535 10.331 1.00 53.78 C
ANISOU 139 CB LEU A 22	5253 7383 7799 -89 -565 -1051 C
ATOM 140 CG LEU A 22	9.539 -2.308 11.786 1.00 55.43 C
ANISOU 140 CG LEU A 22	5484 7602 7974 -188 -393 -1005 C
ATOM 141 CD1 LEU A 22	10.333 -1.048 12.060 1.00 56.18 C
ANISOU 141 CD1 LEU A 22	5655 7678 8014 -100 -364 -953 C
ATOM 142 CD2 LEU A 22	8.337 -2.323 12.714 1.00 57.32 C
ANISOU 142 CD2 LEU A 22	5538 7918 8322 -280 -294 -1108 C
ATOM 143 N ARG A 23	8.778 -2.790 7.031 1.00 49.75 N
ANISOU 143 N ARG A 23	4800 6895 7206 198 -961 -1107 N
ATOM 144 CA ARG A 23	8.808 -3.622 5.795 1.00 56.34 C
ANISOU 144 CA ARG A 23	5682 7751 7974 264 -1112 -1156 C
ATOM 145 C ARG A 23	9.814 -3.086 4.812 1.00 52.25 C
ANISOU 145 C ARG A 23	5320 7263 7270 415 -1167 -1042 C
ATOM 146 O ARG A 23	10.472 -3.867 4.114 1.00 47.78 O

TABLE 3-continued

ANISOU 146 O ARG A 23	4854 5714 6586 447 -1216 -1041 O
ATOM 147 CB ARG A 23	7.446 -3.694 5.064 1.00 61.34 C
ANISOU 147 CB ARG A 23	6156 8439 8713 324 -1274 -1310 C
ATOM 148 CG ARG A 23	6.450 -4.698 5.637 1.00 74.43 C
ANISOU 148 CG ARG A 23	7649 10075 10556 159 -1259 -1448 C
ATOM 149 CD ARG A 23	5.251 -4.971 4.711 1.00 80.04 C
ANISOU 149 CD ARG A 23	8205 10840 11366 221 -1455 -1621 C
ATOM 150 NE ARG A 23	4.820 -6.369 4.853 1.00 90.41 N
ANISOU 150 NE ARG A 23	9441 12098 12812 59 -1477 -1734 N
ATOM 151 CZ ARG A 23	5.184 -7.381 4.052 1.00 93.29 C
ANISOU 151 CZ ARG A 23	9694 12424 13129 73 -1600 -1799 C
ATOM 152 NH1 ARG A 23	5.966 -7.186 2.990 1.00 94.85 N
ANISOU 152 NH1 ARG A 23	10250 12670 13118 252 -1706 -1771 N
ATOM 153 NH2 ARG A 23	4.751 -8.608 4.310 1.00 93.58 N
ANISOU 153 NH2 ARG A 23	9852 12373 13332 -93 -1615 -1897 N
ATOM 154 N VAL A 24	9.920 -1.754 4.754 1.00 49.46 N
ANISOU 154 N VAL A 24	4981 6914 6897 509 -1156 -947 N
ATOM 155 CA VAL A 24	10.883 -1.125 3.856 1.00 46.37 C
ANISOU 155 CA VAL A 24	4731 6552 6336 636 -1187 -800 C
ATOM 156 C VAL A 24	12.280 -1.575 4.276 1.00 44.04 C
ANISOU 156 C VAL A 24	4559 6230 5943 560 -1059 -704 C
ATOM 157 O VAL A 24	13.116 -1.952 3.453 1.00 43.78 O
ANISOU 157 O VAL A 24	4681 6252 5753 624 -1083 -649 O
ATOM 158 CB VAL A 24	10.756 0.404 3.933 1.00 47.32 C
ANISOU 158 CB VAL A 24	4840 6634 6504 718 -1185 -700 C
ATOM 159 CG1 VAL A 24	11.994 1.073 3.364 1.00 47.00 C
ANISOU 159 CG1 VAL A 24	4950 6592 6316 785 -1152 501 C
ATOM 160 CG2 VAL A 24	9.483 0 858 3.161 1.00 47.49 C
ANISOU 160 CG2 VAL A 24	4760 6699 6585 851 -1357 -777 C
ATOM 161 N VAL A 25	12.524 -1.558 5.582 1.00 40.88 N
ANISOU 161 N VAL A 25	4139 5762 5632 430 -925 -695 N
ATOM 162 CA VAL A 25	13.859 -1.929 6.062 1.00 41.30 C
ANISOU 162 CA VAL A 25	4299 5790 5603 365 -817 -605 C
ATOM 163 C VAL A 25	14.041 -3.435 5.925 1.00 40.07 C
ANISOU 163 C VAL A 25	4174 5638 5414 316 -840 -681 C
ATOM 164 O VAL A 25	15.135 -3.887 5.550 1.00 44.59 O
ANISOU 164 O VAL A 25	4846 6228 5868 347 -825 -627 O
ATOM 165 CB VAL A 25	14.073 -1.539 7.547 1.00 40.89 C
ANISOU 165 CB VAL A 25	4222 5679 5637 248 -683 -586 C
ATOM 166 CG1 VAL A 25	15.488 -1.915 7.990 1.00 41.37 C

TABLE 3-continued

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ANISOU 166 CG1 VAL A 25	4387 5722 5609 197 -596 -498 C
ATOM 167 CG2 VAL A 25	13.847 -0.036 7.779 1.00 39.72 C
ANISOU 167 CG2 VAL A 25	4035 5499 5558 298 -670 -547 C
ATOM 168 N ALA A 26	12.981 -4.222 6.224 1.00 42.21 N
ANISOU 168 N ALA A 26	4351 5884 5804 237 -878 -808 N
ATOM 169 CA ALA A 26	13.115 -5.671 6.096 1.00 42.62 C
ANISOU 169 CA ALA A 26	4434 5899 5862 183 -916 -682 C
ATOM 170 C ALA A 26	13.446 -5.975 4.625 1.00 43.94 C
ANISOU 170 C ALA A 26	4667 6135 5895 383 -1046 -924 C
ATOM 171 O ALA A 26	14.376 -6.734 4.326 1.00 46.92 O
ANISOU 171 O ALA A 26	5187 6507 6188 361 -1048 -922 O
ATOM 172 CB ALA A 26	11.855 -6.400 6.606 1.00 42.36 C
ANISOU 172 CB ALA A 26	4272 5818 6006 57 -937 -1000 C
ATOM 173 N ASP A 27	12.777 -5.290 3.708 1.00 45.90 N
ANISOU 173 N ASP A 27	4870 6463 6106 450 -1150 -954 N
ATOM 174 CA ASP A 27	12.956 -5.554 2.245 1.00 45.22 C
ANISOU 174 CA ASP A 27	4842 6481 5859 610 -1286 -1005 C
ATOM 175 C ASP A 27	14.348 -5.193 1.713 1.00 45.46 C
ANISOU 175 C ASP A 27	5002 6588 5684 706 -1222 -867 C
ATOM 176 O ASP A 27	14.937 -5.952 0.892 1.00 45.61 O
ANISOU 176 O ASP A 27	5089 6677 5565 792 -1274 -926 O
ATOM 177 CB ASP A 27	11.916 -4.827 1.397 1.00 46.21 C
ANISOU 177 CB ASP A 27	4895 6690 5972 727 -1424 -1048 C
ATOM 178 CG ASP A 27	10.606 -5.573 1.306 1.00 53.18 C
ANISOU 178 CG ASP A 27	5644 7552 7010 684 -1555 -1250 C
ATOM 179 OD1 ASP A 27	10.608 -6.809 1.454 1.00 53.71 O
ANISOU 179 OD1 ASP A 27	5706 7556 7146 599 -1579 -1368 O
ATOM 180 OD2 ASP A 27	9.555 -4.936 1.067 1.00 55.13 O
ANISOU 180 OD2 ASP A 27	5783 7837 7327 735 -1646 -1295 O
ATOM 181 N TYR A 28	14.832 -4.028 2.134 1.00 45.13 N
ANISOU 181 N TYR A 28	4981 6538 5629 697 -1116 -697 N
ATOM 182 CA TYR A 28	16.185 -3.595 1.787 1.00 40.73 C
ANISOU 182 CA TYR A 28	4520 6042 4912 753 -1028 -544 C
ATOM 183 C TYR A 28	17.212 -4.619 2.283 1.00 43.93 C
ANISOU 183 C TYR A 28	49746415 5302 691 -952 -572 C
ATOM 184 O TYR A 28	18.131 -4.972 1.554 1.00 41.57 O
ANISOU 184 O TYR A 28	4738 6209 4846 778 -943 -556 O
ATOM 185 CB TYR A 28	16.510 -2.185 2.304 1.00 42.18 C
ANISOU 185 CB TR A 28	4704 6182 5141 723 -932 -369 C
ATOM 186 CG TYR A 28	17.901 -1.705 1.956 1.00 41.46 C

TABLE 3-continued

ANISOU 186 CG TYR A 28	4690 6149 4915 756 -835 -201 C
ATOM 187 CD1 TYR A 28	18.126 -0.862 0.860 1.00 44.75 C
ANISOU 187 CD1 TYR A 28	5149 6664 5190 870 -856 -57 C
ATOM 188 CD2 TYR A 28	19.025 -2.070 2.747 1.00 42.22 C
ANISOU 188 CD2 TYR A 28	4807 6207 5027 669 -719 -171 C
ATOM 189 CE1 TYR A 28	19.433 -0.433 0.529 1.00 43.85 C
ANISOU 189 CE1 TYR A 28	5087 6615 4959 881 -747 113 C
ATOM 190 CE2 TYR A 28	20.311 -1.626 2.424 1.00 40.08 C
ANISOU 190 CE2 TYR A 28	4578 6001 4651 691 -626 -25 C
ATOM 191 CZ TYR A 28	20.503 -0.810 1.317 1.00 45.12 C
ANISOU 191 CZ TYR A 28	5246 6741 5158 768 -632 118 C
ATOM 192 OH TYR A 28	21.734 -0.408 1.001 1.00 44.38 O
ANISOU 192 OH TYR A 28	5174 6721 4968 792 -522 272 O
ATOM 193 N ILE A 29	17.061 -5.118 3.507 1.00 40.87 N
ANISOU 193 N ILE A 29	4555 5904 5068 552 -699 -614 N
ATOM 194 CA ILE A 29	18.019 -6.066 4.010 1.00 40.97 C
ANISOU 194 CA ILE A 29	4621 5872 5075 505 -845 -625 C
ATOM 195 C ILE A 29	17.970 -7.390 3.241 1.00 46.14 C
ANISOU 195 C ILE A 29	5302 6538 5691 572 -954 -779 C
ATOM 196 O ILE A 29	19.026 -7.976 2.940 1.00 46 44 O
ANISOU 196 O ILE A 29	5401 6610 5635 635 -938 -788 O
ATOM 197 CB ILE A 29	17.803 -6.330 5.522 1.00 40.88 C
ANISOU 197 CB ILE A 29	4580 5733 5220 343 -771 -617 C
ATOM 198 CG1 ILE A 29	18.006 -5.025 6.311 1.00 38.67 C
ANISOU 198 CG1 ILE A 29	4278 5448 4966 293 -668 -496 C
ATOM 199 CG2 ILE A 29	18.855 -7.307 6.003 1.00 41.88 C
ANISOU 199 CG2 ILE A 29	4773 5809 5332 315 -735 -610 C
ATOM 200 CD1 ILE A 29	17.535 -5.036 7.772 1.00 37.47 C
ANISOU 200 CD1 ILE A 29	4081 5214 4940 150 -596 -501 C
ATOM 201 N LEU A 30	16.760 -7.892 2.983 1.00 44.25 N
ANISOU 201 N LEL A 30	5006 6262 5544 555 -1069 -918 N
ATOM 202 CA LEU A 30	16.557 -9.177 2.297 1.00 49.28 C
ANISOU 202 CA LEU A 30	5658 6879 6189 605 -1199 -1100 C
ATOM 203 C LEU A 30	17.185 -9.121 0.900 1.00 53.86 C
ANISOU 203 C LEU A 30	6295 7630 6538 800 -1259 -1139 C
ATOM 204 O LEU A 30	17.626 -10.162 0.366 1.00 55.53 O
ANISOU 204 O LEU A 30	6549 7847 6701 877 -1329 -1276 O
ATOM 205 CB LEU A 30	15.049 -9.545 2.174 1.00 51.05 C
ANISOU 205 CB LEU A 30	5783 7049 6566 549 -1325 -1247 C
ATOM 206 CG LEU A 30	14.354 -10.172 3.395 1.00 50.90 C

TABLE 3-continued

ANISOU 206 CG LEU A 30	5697 6857 6784 349 -1289 -1265 C
ATOM 207 CD1 LEU A 30	12.800 -10.168 3.263 1.00 49.75 C
ANISOU 207 CD1 LEU A 30	5411 6699 6792 290 -1387 -1385 C
ATOM 208 CD2 LEU A 30	14.898 -11.571 3.648 1.00 54.15 C
ANISOU 208 CD2 LEU A 30	6174 7131 7271 302 -1317 -1331 C
ATOM 209 N GLU A 31	17.196 -7.920 0.307 1.00 47.67 N
ANISOU 209 N GLU A 31	5513 6986 5614 884 -1233 -1021 N
ATOM 210 CA GLU A 31	17.746 -7.730 -1.026 1.00 50.66 C
ANISOU 210 CA GLU A 31	5945 7563 5741 1066 -1269 -1019 C
ATOM 211 C GLU A 31	19.198 -7.394 -1.062 1.00 49.82 C
ANISOU 211 C GLU A 31	5894 7542 5493 1105 -1124 -873 C
ATOM 212 O GLU A 31	19.803 -7.561 -2.103 1.00 49.63 O
ANISOU 212 O GLU A 31	5911 7690 5256 1251 -1134 -900 O
ATOM 213 CB GLU A 31	16.955 -6.718 -1.861 1.00 52.72 C
ANISOU 213 CB GLU A 31	6187 7946 5900 1158 -1344 -962 C
ATOM 214 CG GLU A 31	15.556 -7.226 -2.157 1.00 59.04 C
ANISOU 214 CG GLU A 31	6916 8712 6804 1165 -152 -1160 C
ATOM 215 CD GLU A 31	14.509 -6.134 -2.312 1.00 65.19 C
ANISOU 215 CD GLU A 31	7634 9917 7618 1186 -1590 -1090 C
ATOM 216 OE1 GLU A 31	14.791 -5.085 -2.925 1.00 70.75 O
ANISOU 216 OE1 GLU A 31	8385 10340 8158 1288 -1566 -922 O
ATOM 217 OE2 GLU A 31	13.379 -6.355 -1.841 1.00 70.00 O
ANISOU 217 OE2 GLU A 31	8141 10026 8430 1102 -1668 -1205 O
ATOM 218 N ASN A 32	19.764 -6.927 0.049 1.00 46.51 N
ANISOU 218 N ASN A 32	5467 7021 5183 981 -989 -731 N
ATOM 219 CA ASN A 32	21.134 -6.345 0.023 1.00 49.80 C
ANISOU 219 CA ASN A 32	5910 7527 5483 1005 -845 -567 C
ATOM 220 C ASN A 32	22.045 -6.938 1.058 1.00 42.49 C
ANISOU 220 C ASN A 32	4987 6498 4660 920 -765 -568 C
ATOM 221 O ASN A 32	23.011 -6.305 1.513 1.00 46.19 O
ANISOU 221 O ASN A 32	5447 6982 5121 878 -643 -424 O
ATOM 222 CB ASN A 32	21.062 -4.825 0.214 1.00 47.53 C
ANISOU 222 CB ASS A 32	5606 7242 5211 956 -768 -357 C
ATOM 223 CG ASN A 32	20.366 -4.155 -0.923 1.00 53.03 C
ANISOU 223 CG ASS A 32	6315 8059 5775 1066 -847 -313 C
ATOM 224 OD1 ASN A 32	20.923 -4.046 -1.993 1.00 55.18 O
ANISOU 224 OD1 ASN A 32	6624 8511 5830 1189 -831 -259 O
ATOM 225 ND2 ASN A 32	19.123 -3.729 -0.710 1.00 49.55 N
ANISOU 225 ND2 ASN A 32	5838 7535 5454 1037 -935 -340 N
ATOM 226 N ALA A 33	21.648 -8.116 1.508 1.00 46.62 N

TABLE 3-continued

ANISOU 226 N ALA A 33	5516 6895 5303 882 -845 -723 N
ATOM 227 CA ALA A 33	22.095 -8.660 2.783 1.00 45.21 C
ANISOU 227 CA ALA A 33	5343 6563 5271 769 -797 -709 C
ATOM 228 C ALA A 33	23.591 -8.808 2.847 1.00 44.97 C
ANISOU 228 C ALA A 33	5328 6597 5163 823 -713 -658 C
ATOM 229 O ALA A 33	24.204 -8.507 3.886 1.00 37.52 O
ANISOU 229 O ALA A 33	4375 5590 4292 733 -631 -555 O
ATOM 230 CB ALA A 33	21.417 -9.987 3.104 1.00 48.63 C
ANISOU 220 CB ALA A 33	5790 6843 5846 725 -906 -869 C
ATOM 231 N HIS A 34	24.202 -9.252 1.757 1.00 39.90 N
ANISOU 231 N HIS A 34	4697 6095 4367 977 -734 -740 N
ATOM 232 CA HIS A 34	25.656 -9.313 1.717 1.00 46.43 C
ANISOU 232 CA HIS A 34	5509 7019 5112 1043 -642 -695 C
ATOM 233 C HIS A 34	26.279 -7.971 2.053 1.00 44.81 C
ANISOU 233 C HIS A 34	5261 6881 4863 971 -503 -484 C
ATOM 234 O HIS A 34	27.232 -7.690 2.868 1.00 42.38 O
ANISOU 234 O HIS A 34	4924 6540 4638 918 -431 -417 O
ATOM 235 O3 HIS A 34	26.211 -9.857 0.375 1.00 56.07 C
ANISOU 235 CB HIS A 34	6733 8434 6138 1237 -663 -820 C
ATOM 236 CG HIS A 34	27.131 -9.710 0.254 1.00 62.18 C
ANISOU 236 CG HIS A 34	7457 9355 6815 1305 -538 -755 C
ATOM 237 ND1 HIS A 34	28.581 -10.721 0.522 1.00 61.86 N
ANISOU 237 ND1 HIS A 34	7407 9279 6819 1376 -557 -873 N
ATOM 238 CD2 HIS A 34	28.536 -8.583 -0.050 1.00 59.79 C
ANISOU 238 CD2 HIS A 34	7096 9224 6396 1297 -387 -569 C
ATOM 239 CE1 HIS A 34	29.860 -10.276 0.366 1.00 60.77 C
ANISOU 239 CE1 HIS A 34	7194 9308 6589 1421 -424 -784 C
ATOM 240 NE2 HIS A 34	29.817 -8.977 0.000 1.00 59.58 N
ANISOU 240 NE2 HIS A 34	7013 9284 6342 1363 -316 -595 N
ATOM 241 N ASP A 35	25.791 -6.898 1.434 1.00 41.75 N
ANISOU 241 N ASP A 35	4866 6581 4415 971 -475 -377 N
ATOM 242 CA ASP A 35	26.406 -5.582 1.711 1.00 42.64 C
ANISOU 242 CA ASP A 35	4938 6730 4535 895 -351 -170 C
ATOM 243 C ASP A 35	26.040 -4.957 3.097 1.00 41 15 C
ANISOU 243 C ASP A 35	4735 6356 4543 731 -336 -101 C
ATOM 244 O ASP A 35	26.806 -4.170 3.654 1.00 40.03 O
ANISOU 244 O ASP A 35	4553 6203 4454 657 -247 19 O
ATOM 245 CB ASP A 35	26.116 -4.560 0.608 1.00 48.67 C
ANISOU 245 CB ASP A 35	5706 7.631 5155 950 -324 -42 C
ATOM 246 CG ASP A 35	26.934 -3.281 0.782 1.00 46.96 C

TABLE 3-continued

ANISOU 246 CG ASP A 35	5443 7441 4959 872 -191 179 C
ATOM 247 OD1 ASP A 35	28.191 -3.349 0.785 1.00 54.68 O
ANISOU 247 OD1 ASP A 35	6367 8509 5899 876 -90 225 O
ATOM 248 OD2 ASP A 35	26 329 -2.198 0.987 1.00 53.32 O
ANISOU 248 OD2 ASP A 35	6254 8158 5848 799 -194 301 O
ATOM 249 N VAL A 36	24.881 -5.325 3.636 1 00 36.93 N
ANISOU 249 N VAL A 36	4478 5944 4368 677 -421 -190 N
ATOM 250 CA VAL A 36	24.430 -4.797 4.890 1.00 36.66 C
ANISOU 250 CA VAL A 36	4176 5517 4237 541 -403 -150 C
ATOM 251 C VAL A 36	25.471 -5.159 5.997 1.00 36.13 C
ANISOU 251 C VAL A 36	4100 5397 4230 478 -353 -138 C
ATOM 252 O VAL A 36	25.593 -4.416 6.985 1.00 33.19 O
ANISOU 252 O VAL A 36	3703 4961 3945 379 -307 -73 O
ATOM 253 CB VAL A 36	22.972 -5.227 5.254 1.00 36.96 C
ANISOU 253 CB VAL A 36	4220 5450 4372 491 -489 -254 C
ATOM 254 CG1 VAL A 36	22.598 -4.849 6.718 1.00 35.56 C
ANISOU 254 CG1 VAL A 36	4021 5154 4337 352 -450 -230 C
ATOM 255 CG2 VAL A 36	21.957 -4.592 4.312 1.00 36.36 C
ANISOU 255 CG2 VAL A 36	4132 5425 4258 549 -546 -252 C
ATOM 256 N GLN A 37	26.159 -6.295 5.855 1.00 39.85 N
ANISOU 256 N GLN A 37	4590 5891 4661 546 -378 -215 N
ATOM 257 CA GLN A 37	27.245 -6.717 6.817 1.00 38.48 C
ANISOU 257 CA GLN A 37	4407 5681 4534 517 -351 -204 C
ATOM 258 C GLN A 37	28.230 -5.587 7.067 1.00 39.95 C
ANISOU 258 C GLN A 37	4526 5931 4721 476 -257 -84 C
ATOM 259 O GLN A 37	28 810 -5.437 8.167 1.00 39.29 O
ANISOU 259 O GLN A 37	4422 5798 4709 405 -241 -57 O
ATOM 260 CB GLN A 37	26.084 -7.862 6.223 1.00 42.25 C
ANISOU 260 CB GLN A 37	4892 6216 4945 646 -384 -295 C
ATOM 261 CG GLN A 37	27.349 -9.086 5.777 1.00 44.83 C
ANISOU 261 CG GLN A 37	5278 6477 5280 710 -492 -439 C
ATOM 262 CD GLN A 37	28.303 -10.155 5.243 1.00 45.53 C
ANISOU 262 CD GLN A 37	5369 6614 5317 856 -529 -549 C
ATOM 263 OE1 GLN A 37	28.620 -10.202 4051 1.00 46.35 O
ANISOU 263 OE1 GLN A 37	5450 6872 5290 986 -517 -610 O
ATOM 264 NE2 GLN A 37	28.698 -11.052 6.114 1.00 40.50 N
ANISOU 264 NE2 GLN A 37	4762 5847 4779 846 -582 -580 N
ATOM 265 N PHE A 38	23.423 -4.785 6.036 1.00 41.57 N
ANISOU 265 N PHE A 38	4697 6250 4846 518 -201 -8 N
ATOM 266 CA PHE A 38	29.455 -3.741 6.090 1.00 42.37 C

TABLE 3-continued

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ANISOU 266 CA PHE A 38	4721 6414 4963 474 -106 118 C
ATOM 267 C PHE A 38	28.870 -2.356 6.332 1.00 41.2 C
ANISOU 267 C PHE A 38	4573 6200 4908 376 -82 222 C
ATOM 268 O PHE A 38	29.623 -1.398 6.256 1.00 38.70 O
ANISOU 268 O PHE A 38	4164 5903 4617 328 -9 387 O
ATOM 269 CB PHE A 38	30.362 -3.768 4.839 1.00 40.21 C
ANISOU 269 CB PHE A 38	4398 6329 4550 577 -33 163 C
ATOM 270 CG PHE A 38	30.824 -5.150 4.467 1.00 43.62 C
ANISOU 270 CG PHE A 38	4840 6836 4899 706 -71 23 C
ATOM 271 CD1 PHE A 38	31.780 -5.818 5.229 1.00 40.97 C
ANISOU 271 CD1 PHE A 38	4464 6479 4624 716 -81 -36 C
ATOM 272 CD2 PHE A 38	30.318 -5.786 3.324 1.00 43.50 C
ANISOU 272 CD2 PHE A 38	4872 6913 4744 834 -111 -63 C
ATOM 273 CE1 PHE A 38	32.200 -7.117 4 897 1.00 38.72 C
ANISOU 273 CE1 PHE A 38	4190 6237 4286 852 -132 -176 C
ATOM 274 CE2 PHE A 38	30.725 -7.069 2.988 1.00 47.43 C
ANISOU 274 CE2 PHE A 38	5378 7459 5184 964 -161 -221 C
ATOM 275 CZ PHE A 38	31.661 -7.746 3.792 1.00 48.72 C
ANISOU 275 CZ PHE A 38	5505 7574 5433 974 -171 -276 C
ATOM 276 N GLN A 39	27.559 -2.259 6.686 1.00 34.59 N
ANISOU 276 N GLN A 39	3777 5249 4117 342 -144 176 N
ATOM 277 CA GLN A 39	26.935 -0.950 6.883 1.00 32.55 C
ANISOU 277 CA GLN A 39	3506 4912 3951 274 -135 252 C
ATOM 278 C GLN A 39	26.790 -0.709 8.3638 1.00 34.18 C
ANISOU 278 C GLN A 39	3699 5001 4285 172 -146 197 C
ATOM 279 O GLN A 39	26.562 -1.657 9.105 1.00 35.04 O
ANISOU 279 O GLN A 39	3837 5085 4393 158 -179 102 O
ATOM 280 CB GLN A 39	25.511 -0.912 6.259 1.00 34.84 C
ANISOU 280 CB GLN A 39	3837 5183 4217 323 -202 220 C
ATOM 281 CG GLN A 39	25.550 -0.947 4.724 1.00 39.65 C
ANISOU 281 CG GLN A 39	4466 5926 4673 438 -204 282 C
ATOM 282 CD GLN A 39	24.177 -0.751 4.066 1.00 41.60 C
ANISOU 282 CD GLN A 39	4745 6164 4898 496 -291 257 C
ATOM 283 OE1 GLN A 39	23.262 -0.174 4.657 1.00 41.02 O
ANISOU 283 OE1 GLN A 39	4658 5977 4949 446 -328 237 O
ATOM 284 NE2 GLN A 39	24.054 -1.211 2.830 1.00 40.35 N
ANISOU 284 NE2 GLN A 39	4617 6140 4576 614 -327 247 N
ATOM 285 N THR A 40	26.870 0.553 8.738 1.00 34.94 N
ANISOU 285 N THR A 40	3758 5029 4487 105 -122 259 N
ATOM 286 CA THR A 40	26.604 1.015 10.080 1.00 36.19 C

TABLE 3-continued

ANISOU 286 CA THR A 40	3902 5090 4760 21 -136 190 C
ATOM 287 C THR A 40	25.117 1.259 10.255 1.00 34.18 C
ANISOU 287 C THR A 40	3667 4770 4549 25 -176 125 C
ATOM 288 O THR A 40	24.356 1.251 9.276 1.00 36.16 O
ANISOU 288 O THR A 40	3937 5039 4764 89 -203 151 O
ATOM 289 CB THR A 40	27.368 2.319 10.391 1.00 34.41 C
ANISOU 289 CB THR A 40	3616 4802 4656 -47 -108 257 C
ATOM 290 OG1 THR A 40	26.833 3.405 9.604 1.00 33.29 O
ANISOU 290 OG1 THR A 40	3473 4596 4578 -34 -109 352 O
ATOM 291 CG2 THR A 40	28.854 2.104 10.115 1.00 30.81 C
ANISOU 291 CG2 THR A 40	3109 4429 4168 -55 -60 324 C
ATOM 292 N ILE A 41	24.674 1.471 11.496 1.00 34.84 N
ANISOU 292 N ILE A 41	3738 4797 4702 -36 -182 32 N
ATOM 293 CA ILE A 41	23.228 1.755 11.667 1.00 36.82 C
ANISOU 293 CA ILE A 41	3981 5004 5006 -28 -208 -42 C
ATOM 294 C ILE A 41	22.906 3.010 10.863 1.00 38.44 C
ANISOU 294 C ILE A 41	4166 5143 5298 12 -231 30 C
ATOM 295 O ILE A 41	21.795 3.108 10.261 1.00 37.24 O
ANISOU 295 O ILE A 41	4011 4983 5156 70 -273 15 O
ATOM 296 CB ILE A 41	22.834 1.861 18.147 1.00 36.95 C
ANISOU 296 CB ILE A 41	3975 4999 5064 -92 -193 -158 C
ATOM 297 CG1 ILE A 41	21.323 1.995 13.276 1.00 40.02 C
ANISOU 297 CG1 ILE A 41	4332 5374 5499 -79 -205 -245 C
ATOM 298 CG2 ILE A 41	23.722 2.883 13.871 1.00 32.61 C
ANISOU 298 CG2 ILE A 41	3422 4425 4619 -136 -186 -166 C
ATOM 299 CD1 ILE A 41	20.904 1.832 14.723 1.00 43.04 C
ANISOU 299 CD2 ILE A 42	4692 5786 5874 -140 -166 -358 C
ATOM 300 N THR A 42	23.895 3.932 10.800 1.00 35.28 N
ANISOU 300 N THR A 42	3747 4691 4965 -17 -211 118 N
ATOM 301 CA THR A 42	23.732 5.285 10.150 1.00 36.89 C
ANISOU 301 CA THR A 42	3938 4790 5287 3 -234 218 C
ATOM 302 C THR A 42	23.549 5.083 8.644 1.00 35.03 C
ANISOU 302 C THR A 42	3741 4623 4946 87 -245 353 C
ATOM 303 C THR A 42	22.665 5.674 8.031 1.00 35.95 O
ANISOU 303 O THR A 42	3869 4690 5101 150 -298 393 O
ATOM 304 CB THR A 42	24.914 6.266 10.454 1.00 36.99 C
ANISOU 304 CB THR A 42	3915 4717 5423 -74 -208 291 C
ATOM 305 OG1 THR A 42	25.331 6.131 11.821 1.00 39.80 O
ANISOU 305 OG1 THR A 42	4240 5066 5816 -141 -204 152 O
ATOM 306 CG2 THR A 42	24.571 7.726 10.194 1.00 39.76 C

TABLE 3-continued

ANISOU 306 CG2 THR A 42	4253 4899 5955 -72 -249 359 C
ATOM 307 N ASP A 43	24.282 4.146 8.071 1.00 35.09 N
ANISOU 307 N ASP A 43	3769 4758 4804 105 -206 400 N
ATOM 308 CA ASP A 43	24.066 3.794 6.623 1.00 36.36 C
ANISOU 308 CA ASP A 43	3971 5026 4818 204 -220 494 C
ATOM 309 C ASP A 43	22.738 3.134 6.354 1.00 38.06 C
ANISOU 309 C ASP A 43	4208 5272 4981 277 -298 383 C
ATOM 310 O ASP A 43	22.029 6026 5.404 1.00 43.89 O
ANISOU 310 O ASP A 43	4967 6026 5683 361 -356 442 O
ATOM 311 CB ASP A 43	25.092 2.741 6.153 1.00 37.67 C
ANISOU 311 CB ASP A 43	4144 5342 4827 227 -166 510 C
ATOM 312 CG ASP A 43	26.494 3.314 5.995 1.00 39.98 C
ANISOU 312 CG ASP A 43	4394 5659 5137 173 -79 650 C
ATOM 313 OD1 ASP A 43	26.657 4.441 5.462 1.00 41.53 O
ANISOU 313 OD1 ASP A 43	4582 5807 5391 155 -56 812 O
ATOM 314 OD2 ASP A 43	27.441 2.581 6.364 1.00 43.89 O
ANISOU 314 OD2 ASP A 43	4860 6226 5592 152 -37 602 O
ATOM 315 N LEU A 44	22.403 2.117 7.143 1.00 39.04 N
ANISOU 315 N LEU A 44	4325 5409 5100 246 -306 230 N
ATOM 316 CA LEU A 44	21.123 1.373 6.901 1.00 40.82 C
ANISOU 316 CA LEU A 44	4550 5660 5298 295 -379 114 C
ATOM 317 C LEU A 44	19.921 2.327 6.955 1.00 43.83 C
ANISOU 317 C LEU A 44	4893 5963 5796 320 -436 91 C
ATOM 318 O LEU A 44	18.985 2.230 6.132 1.00 44.18 O
ANISOU 318 O LEU A 44	4933 6044 5809 403 -518 68 O
ATOM 319 CB LEU A 44	20.937 0.301 7.954 1.00 39.47 C
ANISOU 319 CB LEU A 44	4370 5480 5145 224 -364 -18 C
ATOM 320 CG LEU A 44	19.737 -0.605 7.708 1.00 45.48 C
ANISOU 320 CG LEU A 44	5118 6262 5901 246 -431 -133 C
ATOM 321 CD1 LEU A 44	19.705 -1.107 6.280 1.00 41.22 C
ANISOU 321 CD1 LEU A 44	4612 5809 5239 354 -499 -121 C
ATOM 322 CD2 LEU A 44	19.730 -1.750 8.694 1.00 39.47 C
ANISOU 322 CD2 LEU A 44	4360 5482 5154 161 -405 -218 C
ATOM 323 N ALA A 45	19.973 3.258 7.911 1.00 43.67 N
ANISOU 323 N ALA A 45	4839 5840 5913 260 -406 81 N
ATOM 324 CA ALA A 45	18.980 4.303 8.043 1.00 45.17 C
ANISOU 324 CA ALA A 45	4986 5938 6240 296 -459 50 C
ATOM 325 C ALA A 45	18.911 5.2056 6.820 1.00 45.80 C
ANISOU 325 C ALA A 45	5097 5987 6317 390 -520 205 C
ATOM 326 O ALA A 45	17.806 5.427 6.252 1.00 45.33 O

TABLE 3-continued

ANISOU 326 O ALA A 45	5018 5927 6280 482 -611 180 O
ATOM 327 CB ALA A 45	19.176 5.107 9.324 1.00 41.04 C
ANISOU 327 CB ALA A 45	4423 5311 5858 224 -418 -15 C
ATOM 328 N ARG A 46	20.058 5.720 5.396 1.00 44.63 N
ANISOU 328 N ARG A 46	4992 5822 6144 369 -473 373 N
ATOM 329 CA ARG A 46	20.108 6.535 5.185 1.00 48.79 C
ANISOU 329 CA ARG A 46	5562 6333 6642 448 -515 569 C
ATOM 330 C ARG A 46	19.662 5.802 3.935 1.00 48.49 C
ANISOU 330 C ARG A 46	5565 6453 6407 560 -572 599 C
ATOM 331 O ARG A 46	18.871 6.321 3.147 1.00 48.11 O
ANISOU 331 O ARG A 46	5534 6396 6351 664 -668 662 O
ATOM 332 CB ARG A 46	21.551 6.995 4.907 1.00 53.91 C
ANISOU 332 CB ARG A 46	6237 6976 7272 382 -425 758 C
ATOM 333 CG ARG A 46	22.129 8.004 5.863 1.00 55.16 C
ANISOU 333 CG ARG A 46	6359 6959 7642 278 -391 771 C
ATOM 334 CD ARG A 46	23.608 8.211 5.499 1.00 61.83 C
ANISOU 334 CD ARG A 46	7204 7834 8453 199 -294 950 C
ATOM 335 NE ARG A 46	23.822 9.304 4.526 1.00 61.58 N
ANISOU 335 NE ARG A 46	7206 7726 8465 214 -297 1205 N
ATOM 336 CZ ARG A 46	24.080 9.167 3.223 1.00 64.41 C
ANISOU 336 CZ ARG A 46	7613 8224 8636 275 -266 1403 C
ATOM 337 NH1 ARG A 46	24.183 7.978 2.618 1.00 66.99 N
ANISOU 337 NH1 ARG A 46	7959 8781 8715 343 -235 1358 N
ATOM 338 NH2 ARG A 46	24.260 10.249 2.514 1.00 62.68 N
ANISOU 338 NH2 ARG A 46	7425 7911 8478 271 -266 1651 N
ATOM 339 N ASN A 47	20.227 4.621 3.701 1.00 45.22 N
ANISOU 339 N ASN A 47	5168 6183 5830 551 -525 555 N
ATOM 340 CA ASN A 47	19.899 3.892 2.485 1.00 46.66 C
ANISOU 340 CA ASN A 47	5390 6525 5813 665 -586 557 C
ATOM 341 C ASN A 47	18.414 3.501 2.381 1.00 49.08 C
ANISOU 341 C ASN A 47	5663 6638 6149 736 -714 395 C
ATOM 342 O ASN A 47	17.917 3.236 1.238 1.00 52.38 O
ANISOU 342 O ASN A 47	6108 7367 6428 853 -806 402 O
ATOM 343 CB ASN A 47	20.741 2.642 2.406 1.00 47.08 C
ANISOU 343 CB ASN A 47	5457 6707 5723 648 -522 490 C
ATOM 344 CG ASN A 47	22.208 2.962 2.328 1.00 48.03 C
ANISOU 344 CG ASN A 47	5588 6862 5798 597 -400 644 C
ATOM 345 OD1 ASN A 47	22.587 4.050 1.864 1.00 43.26 O
ANISOU 345 OD1 ASN A 47	4998 6231 5207 596 -367 842 O
ATOM 346 ND2 ASN A 47	23.038 2.046 2.797 1.00 43.71 N

TABLE 3-continued

ANISOU 346 ND2 ASN A 47	5026 6365 5215 550 -334 562 N
ATOM 347 N THR A 48	17.737 3.415 3.524 1.00 43.99 N
ANISOU 347 N THR A 48	4949 6093 5672 664 -717 242 N
ATOM 348 CA THR 4 48	16.336 3.068 3.509 1.00 46.99 C
ANISOU 348 CA THR A 48	5265 6482 6108 712 -824 85 C
ATOM 349 C THR A 48	15.430 4.301 3.680 1.00 45.39 C
ANISOU 349 C THR A 48	5010 6168 6068 763 -892 98 C
ATOM 350 O THR A 48	14.196 4.171 3.614 1.00 45.99 O
ANISOU 350 O THR A 48	5009 6257 6209 816 -990 -30 O
ATOM 351 CB THR A 48	15.995 2.098 4.647 1.00 42.76 C
ANISOU 351 CB THR A 48	4666 5929 5653 602 -778 -98 C
ATOM 352 OG1 THR A 48	16.378 2.703 5.885 1.00 39.24 O
ANISOU 352 OG1 THR A 48	4196 5381 5332 504 -683 -95 O
ATOM 353 CG2 THR A 48	16.642 0.692 4.409 1.00 42.62 C
ANISOU 353 CG2 THR A 48	4694 6000 5500 574 -751 -145 C
ATOM 354 N GLN A 49	16.041 5.465 3.900 1.00 48.72 N
ANISOU 354 N GLN A 49	5462 6475 6573 746 -849 241 N
ATOM 355 CA GLN A 49	15.319 6.701 4.257 1.00 53.62 C
ANISOU 355 CA GLN A 49	6036 6947 7390 790 -910 238 C
ATOM 356 C GLN A 49	14.355 6.500 5.450 1.00 52.27 C
ANISOU 356 C GLN A 49	5746 6742 7372 746 -902 2 C
ATOM 357 O GLN A 49	13.170 6.870 5.384 1.00 54.02 O
ANISOU 357 O GLN A 49	5886 6940 7698 833 -999 -91 O
ATOM 358 CB GLN A 49	14.513 7.213 3.057 1.00 58.34 C
ANISOU 358 CB GLN A 49	6652 7563 7953 951 -1059 325 C
ATOM 359 CG GLN A 49	15.296 7.882 1.925 1.00 70.89 C
ANISOU 359 CG GLN A 49	8356 9155 9423 1011 -1072 603 C
ATOM 360 CD GLN A 49	14.441 7.949 0.661 1.00 80.36 C
ANISOU 360 CD GLN A 49	9582 10448 10504 1183 -1231 663 C
ATOM 361 OE1 GLN A 49	14.248 6.939 -0.021 1.00 89.83 O
ANISOU 361 OE1 GLN A 49	10790 11829 11511 1232 -1272 594 O
ATOM 362 NE2 GLN A 49	13.894 9.127 0.360 1.00 88.39 N
ANISOU 362 NE2 GLN A 49	10609 11334 11642 1285 -1340 775 N
ATOM 363 N THR A 50	14.845 5.915 6.539 1.00 45.32 N
ANISOU 363 N THR A 50	4846 5874 6499 617 -787 -91 N
ATOM 364 CA THR A 50	14.050 5.708 7.748 1.00 43.84 C
ANISOU 364 CA THH A 50	4552 5683 6424 561 -748 -291 C
ATOM 365 C THR A 50	14.929 6.228 8.880 1.00 46.84 C
ANISOU 365 C THR A 50	4949 5981 6568 467 -648 -295 C
ATOM 366 O THR A 50	16.120 6.465 8.686 1.00 57.26 O

TABLE 3-continued

ANISOU 366 O THR A 50	6351 7264 8143 431 -611 -158 O
ATOM 367 CB THR A 50	13.621 4.201 7.914 1.00 45.44 C
ANISOU 367 CB THR A 50	4715 6012 6537 495 -720 -406 C
ATOM 368 OG1 THR A 50	14.766 3.357 8.167 1.00 46.98 O
ANISOU 368 OG1 THR A 50	4991 6245 6614 399 -632 -353 O
ATOM 369 OG2 THR A 50	12.967 3.662 6.650 1.00 44.81 C
ANISOU 369 OG2 THR A 50	4631 6012 6382 589 -839 -404 C
ATOM 370 N SER A 51	14.405 6.386 10.072 1.00 51.12 N
ANISOU 370 N SER A 51	5406 6512 7505 426 -601 -459 N
ATOM 371 CA SER A 51	15.267 6.781 11.187 1.00 51.23 C
ANISOU 371 CA SER A 51	5438 6474 7552 342 -518 -488 C
ATOM 372 C SER A 51	16.129 5.590 11.683 1.00 51.03 C
ANISOU 372 C SER A 51	5465 6549 7376 228 -426 -470 C
ATOM 373 O SER A 51	15.882 4.428 11.359 1.00 44.63 O
ANISOU 373 O SER A 51	4661 5835 6462 20 -417 -469 O
ATOM 374 CB SER A 51	14.402 7.258 12.303 1.00 50.13 C
ANISOU 374 CB SER A 51	5192 6328 7527 350 -493 -686 C
ATOM 375 OG SER A 51	13.577 6.178 12.627 1.00 53.50 O
ANISOU 375 OG SER A 51	5550 6895 7884 309 -446 -785 O
ATOM 376 N GLU A 52	17.125 5.880 12.503 1.00 46.44 N
ANISOU 376 N GLU A 52	4914 5934 6796 159 -371 -468 N
ATOM 377 CA GLU A 52	17.815 4.813 13.172 1.00 45.04 C
ANISOU 377 CA GLU A 52	4774 5849 6491 67 -298 -473 C
ATOM 378 C GLU A 52	16.890 4.136 14.165 1.00 42.54 C
ANISOU 378 C GLU A 52	4394 -5625 6143 21 -241 -614 C
ATOM 379 O GLU A 52	17.047 2.958 14.454 1.00 44.75 O
ANISOU 379 O GLU A 52	4704 5986 6314 -45 -196 -599 O
ATOM 380 CB GLU A 52	19.041 5.341 13.892 1.00 45.10 C
ANISOU 380 CB GLU A 52	4812 5809 6514 12 -270 -458 C
ATOM 381 CG GLU A 52	19.994 6.056 12.973 1.00 50.45 C
ANISOU 381 CG GLU A 52	5531 6395 7242 32 -305 -304 C
ATOM 382 CD GLU A 52	21.295 6.439 13.673 1.00 54.72 C
ANISOU 382 CD GLU A 52	6063 6900 7810 -40 -281 -294 C
ATOM 383 OE1 GLU A 52	21.502 6.036 14.845 1.00 53.04 O
ANISOU 383 OE1 GLU A 52	5660 6750 7544 -92 -247 -406 O
ATOM 384 OE2 GLU A 52	22.117 7.123 13.023 1.00 58.16 O
ANISOU 384 OE2 GLU A 52	6532 7253 8315 -47 -298 -166 O
ATOM 385 N ALA A 53	15.938 4.670 14.696 1.00 41.68 N
ANISOU 385 N ALA A 53	4197 5506 6135 57 -240 -747 N
ATOM 386 CA ALA A 53	14.939 4.219 15.609 1.00 50.52 C

TABLE 3-continued

ANISOU 386 CA ALA A 53	5232 6745 7219 8 -163 -876 C
ATOM 387 C ALA A 53	14.134 3.084 14.92 1.00 46.94 C
ANISOU 387 C ALA A 53	4748 6354 6733 -12 -172 -848 C
ATOM 388 O ALA A 53	13.753 2.079 15.550 1.00 44.7 O
ANISOU 388 O ALA A 53	4445 6170 6392 -102 -98 -875 O
ATOM 389 CB ALA A 53	13.972 5.248 16.168 1.00 46.28 C
ANISOU 389 CB ALA A 53	4831 6456 7057 73 -160 -1044 C
ATOM 390 N THR A 54	13.810 3.310 13.655 1.00 48.56 N
ANISOU 390 N THR A 54	4950 6506 6994 74 -272 -798 N
ATOM 391 CA THR A 54	13.043 2.333 2.333 12.873 1.00 47.18 C
ANISOU 391 CA THR A 54	4740 5381 6804 72 -315 -797 C
ATOM 392 C THR A 54	13.898 1.055 12.699 1.00 45.63 C
ANISOU 392 C THR A 54	4648 6207 6463 -5 -294 -702 C
ATOM 393 O THR A 54	13.425 -0.059 12.933 1.00 42.99 O
ANISOU 393 O THR A 54	4285 5919 6129 -84 -265 -732 O
ATOM 394 CB THR A 54	12.605 2.892 11.510 1.00 48.29 C
ANISOU 394 CB THR A 54	4869 6476 7002 202 -446 -764 C
ATOM 395 OG1 THR A 54	11.84 4.084 11.686 1.00 54.69 O
ANISOU 395 OG1 THR A 54	5587 7245 7949 289 -481 -849 O
ATOM 396 CG2 THR A 54	11.723 1.859 10.770 1.00 46.86 C
ANISOU 396 CG2 THR A 54	4632 6360 6812 201 -508 -805 C
ATOM 397 N VAL A 55	15.173 1.246 12.362 1.00 39.75 N
ANISOU 397 N VAL A 55	4015 5420 5670 16 -306 -591 N
ATOM 398 CA VAL A 55	16.080 0.124 12.117 1.00 41.43 C
ANISOU 398 CA VAL A 55	4321 5648 5772 -26 -299 -512 C
ATOM 399 C VAL A 55	16.210 -0.671 13.413 1.00 40.01 C
ANISOU 399 C VAL A 55	4148 5502 5552 -141 -210 -538 C
ATOM 400 O VAL A 55	16.116 -1.899 13.394 1.00 40.78 O
ANISOU 400 O VAL A 55	4270 5610 5614 -195 -209 -527 O
ATOM 401 CB VAL A 55	17.463 0.661 11.647 1.00 41.50 C
ANISOU 401 CB VAL A 55	4416 5624 5728 17 -309 -396 C
ATOM 402 CG1 VAL A 55	18.478 -0.453 11.456 1.00 39.61 C
ANISOU 402 CG1 VAL A 55	4258 5411 5380 -7 -299 -334 C
ATOM 403 CG2 VAL A 55	17.319 1.451 10.351 1.00 46.94 C
ANISOU 403 CG2 VAL A 55	5109 6289 6438 126 -387 -333 C
ATOM 404 N VAL A 56	16.399 0.052 14.536 1.00 39.35 N
ANISOU 404 N VAL A 56	4045 5431 5474 -172 -145 -575 N
ATOM 405 CA VAL A 56	16.532 -0.586 15.837 1.00 41.03 C
ANISOU 405 CA VAL A 56	4271 5701 5616 -270 -59 -588 C
ATOM 406 C VAL A 56	15.267 -1.388 16.164 1.00 42.41 C

TABLE 3-continued

ANISOU 406 C VAL A 56	4367 5931 5817 -343 -11 -639 C
ATOM 407 O VAL A 56	15.356 -2.550 16.606 1.00 41.25 O
ANISOU 407 O VAL A 56	4262 5798 5612 -432 26 -584 O
ATOM 408 CB VAL A 56	16.808 0.438 16.980 1.00 43.27 C
ANISOU 408 CB VAL A 56	4534 6015 5891 -272 -7 -657 C
ATOM 409 CG1 VAL A 56	16.804 -0.264 18.304 1.00 41.87 C
ANISOU 409 CG1 VAL A 56	4372 5933 5605 -364 82 -665 C
ATOM 410 CG2 VAL A 56	18.160 1.157 16.822 1.00 42.91 C
ANISOU 410 CG2 VAL A 56	4556 5908 5841 -232 -51 -606 C
ATOM 411 N ARG A 57	14.090 -0.782 15.930 1.00 42.94 N
ANISOU 411 N ARG A 57	4312 6018 5985 -307 -17 -738 N
ATOM 412 CA ARG A 57	12.850 -1.458 16.227 1.00 45.14 C
ANISOU 412 CA ARG A 57	4480 6360 6312 -385 36 -796 C
ATOM 413 C ARG A 57	12.727 -2.734 15.419 1.00 39.67 C
ANISOU 413 C ARG A 57	3818 5618 5636 -431 -24 -739 C
ATOM 414 O ARG A 57	12.346 -3.773 15.964 1.00 43.90 O
ANISOU 414 O ARG A 57	4337 6173 6170 -552 37 -713 O
ATOM 415 CB ARG A 57	11.619 -0.530 16.057 1.00 49.08 C
ANISOU 415 CB ARG A 57	4819 6895 6933 -319 25 -932 C
ATOM 416 CG ARG A 57	10.384 -1.007 16.812 1.00 58.34 C
ANISOU 416 CG ARG A 57	5840 8179 8149 -416 131 -1012 C
ATOM 417 CD ARG A 57	9.190 -0.060 16.633 1.00 72.05 C
ANISOU 417 CD ARG A 57	7396 9960 10018 -328 113 -1167 C
ATOM 418 NE ARG A 57	9 578 1.119 15.853 1.00 81.97 N
ANISOU 418 NE ARG A 57	8696 11120 11327 -159 -8 -1182 N
ATOM 419 CZ ARG A 57	8.858 1.664 14.872 1.00 83.53 C
ANISOU 419 CZ ARG A 57	8813 11279 11645 -55 -126 -1239 C
ATOM 420 NH1 ARG A 57	7.666 1.156 14.555 1.00 86.02 N
ANISOU 420 NH1 ARG A 57	8976 11656 12052 -77 -144 -1317 N
ATOM 421 NH2 ARG A 57	9.337 2.727 14.215 1.00 78.00 N
ANISOU 421 NH2 ARG A 57	8180 10475 10980 79 -230 -1210 N
ATOM 422 N LEU A 58	13.078 -2.703 14.136 1.00 43.67 N
ANISOU 422 N LEU A 58	4377 6060 6156 -337 -144 -716 N
ATOM 423 CA LEU A 58	13.112 -3.945 13-354 1.00 41.71 C
ANISOU 423 C ALEU A 58	4172 5763 5912 -363 -216 -689 C
ATOM 424 C LEU A 58	14.026 -4.997 14.055 1.00 43.03 C
ANISOU 424 C LEU A 58	4454 5894 6000 -452 -167 -592 C
ATOM 425 O LEU A 58	13.612 -6.138 14.353 1.00 42.00 O
ANISOU 425 O LEU A 58	4316 5731 5910 -560 -149 -577 O
ATOM 426 CB LEU A 58	13.571 -3.645 11.906 1.00 42.99 C

TABLE 3-continued

ANISOU 426 CB LEU A 58	4391 5895 6047 -224 -342 -677 C
ATOM 427 CG LEU A 58	13.785 -4.648 10.989 1.00 44.91 C
ANISOU 427 CG LEU A 58	4694 6098 6272 -214 -434 -677 C
ATOM 428 CD1 LEU A 58	12.470 -5.550 10.766 1.00 46.33 C
ANISOU 428 CD1 LEU A 58	4758 6273 -275 -482 -779 C
ATOM 429 CD2 LEU A 58	14.343 -4.503 9.631 1.00 41.98 C
ANISOU 429 CD2 LEU A 58	4365 5741 5825 -67 -636 -560 C
ATOM 430 N CYS A 59	15.296 -4.609 14.310 1.00 39.52 N
ANISOU 430 N CYS A 59	4115 5445 5456 -406 -153 -521 N
ATOM 431 CA CYS A 59	16.221 -5.451 15.016 1.00 38.24 C
ANISOU 431 CA CYS A 59	4057 5257 5216 -464 -122 -433 C
ATOM 432 C CYS A 59	15.604 -6.042 16.279 1.00 37.83 C
ANISOU 432 C CYS A 59	3976 5237 5161 -601 -22 -407 C
ATOM 433 O CYS A 59	15.639 -7.274 16.453 1.00 39.95 O
ANISOU 433 O CYS A 59	4297 5442 5440 -679 -29 -341 O
ATOM 434 CB CYS A 59	17.573 -4.691 15.307 1.00 31.78 C
ANISOU 434 CB CYS A 59	3312 4456 4306 -400 -114 -384 C
ATOM 435 SG CYS A 59	18.316 -4.365 13.676 1.00 38.09 S
ANISOU 435 SG CYS A 59	4148 5223 5100 -263 -215 -372 S
ATOM 436 N ARG A 60	15.044 -5.218 17.162 1.00 37.75 N
ANISOU 436 N ARG A 60	3886 5323 5135 -631 72 -454 N
ATOM 437 CA ARG A 60	14.535 -5.794 18.402 1.00 43.59 C
ANISOU 437 CA ARG A 60	4602 6127 5832 -762 188 -410 C
ATOM 438 C ARG A 60	13.264 -6.636 18.197 1.00 47.77 C
ANISOU 438 C ARG A 60	5029 6639 6482 -872 213 -423 C
ATOM 439 O ARG A 60	13.094 -7.656 18.860 1.00 47.28 O
ANISOU 439 O ARG A 60	4994 6562 6408 -1000 273 -323 O
ATOM 440 CB ARG A 60	14.314 -4.721 19.450 1.00 44.11 C
ANISOU 440 CB ARG A 60	4606 6329 5826 -755 290 -478 C
ATOM 441 CG ARG A 60	15.598 -4.019 19 790 1.00 47.38 C
ANISOU 441 CG ARG A 60	5117 6748 6136 -674 258 -467 C
ATOM 442 CD ARG A 60	15.399 -3.025 20.914 1.00 58.02 C
ANISOU 442 CD ARG A 60	6409 8228 7409 -865 346 -561 C
ATOM 443 NE ARG A 60	16.709 -2.419 21.194 1.00 63.91 N
ANISOU 443 NE ARG A 60	7244 8959 8078 -597 292 -558 N
ATOM 444 CZ ARG A 60	16.925 -1.183 21.657 1.00 60.67 C
ANISOU 444 CZ ARG A 60	6799 8593 7661 -533 293 -676 C
ATOM 445 NH1 ARG A 60	15.915 -0.368 21.897 1.00 58.59 N
ANISOU 445 NH1 ARG A 60	6416 8392 7454 -509 346 -816 N
ATOM 446 NH2 ARG A 60	18.177 -0.762 21.877 1.00 52.18 N

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TABLE 3-continued

ANISOU 446 NH2 ARG A 60	5799 7492 6536 -491 231 -868 N
ATOM 447 N ASP A 61	12.434 -6.257 17 221 1.00 49.01 N
ANISOU 447 N ASP A 61	5073 6785 6764 -821 150 -534 N
ATOM 448 CA ASP A 61	11.227 -7.018 16.923 1.00 52.46 C
ANISOU 448 CA A3P A 61	5388 7202 7342 -922 151 -571 C
ATOM 449 C ASP A 61	11.627 -8.407 16.526 1.00 57.44 C
ANISOU 449 C ASP A 61	6120 7690 8014 -986 76 -486 C
ATOM 450 O ASP A 61	10.846 -9.374 16.690 1.00 59.21 O
ANISOU 450 O ASP A 61	6279 7868 8350 -1129 102 -462 O
ATOM 451 CB ASP A 61	10.402 -6.383 15.797 1.00 54.41 C
ANISOU 451 CB ASP A 61	5507 7456 7709 -825 53 -714 C
ATOM 452 CG ASP A 61	9.570 -5.217 16.275 1.00 56.33 C
ANISOU 452 CG ASP A 61	5598 7828 7978 -790 131 -819 C
ATOM 453 OD1 ADP A 61	9.607 -4.900 17.478 1.00 52.76 O
ANISOU 453 OD1 ASP A 61	5132 7472 7442 -841 270 -800 O
ATOM 454 OD2 ASP A 61	8.883 -4.610 15.438 1.00 59.70 O
ANISOU 454 OD2 ASP A 61	5918 8263 8503 -695 45 -930 O
ATOM 455 N MET A 62	12.839 -8.526 16.006 1.00 52.50 N
ANISOU 455 N MET A 62	5644 6989 7314 -884 -16 -445 N
ATOM 456 CA MET A 62	13.270 -9.815 15.480 1.00 51.82 C
ANISOU 456 CA MET A 62	5655 6755 7280 -907 -113 -399 C
ATOM 457 C MET A 62	13.980 -10.571 16.548 1.00 52.66 C
ANISOU 457 C MET A 62	5881 6816 7313 -991 -51 -245 C
ATOM 458 O MET A 62	14.495 -11.643 16.271 1.00 47.02 O
ANISOU 458 O MET A 62	5265 5961 6640 -999 -132 -194 O
ATOM 459 CB MET A 62	14.179 -9.663 14.297 1.00 56.89 C
ANISOU 459 CB MET A 62	6382 7354 7880 -743 -243 -446 C
ATOM 460 CG MET A 62	13.602 -8.747 13.241 1.00 59.05 C
ANISOU 460 CG MET A 62	6560 7693 8184 -635 -309 -570 C
ATOM 461 SD MET A 62	13.988 -9.733 11.830 1.00 77.50 S
ANISOU 461 SD MET A 62	8968 9927 10552 -547 -478 -634 S
ATOM 462 CE MET A 62	12.669 -10.948 12.042 1.00 50.44 C
ANISOU 462 CE MET A 62	5438 6399 7327 -724 -498 -585 C
ATOM 463 N GLY A 63	13.983 -10.022 17.769 1.00 47.98 N
ANISOU 463 N GLY A 63	5278 6343 6609 -1042 83 -181 N
ATOM 464 CA GLY A 63	14.541 -10.726 18.910 1.00 49.98 C
ANISOU 464 CA GLY A 63	5643 6581 6768 -1127 146 -17 C
ATOM 465 C GLY A 63	16.030 -10 462 19.127 1.00 50.07 C
ANISOU 465 C GLY A 63	5795 6596 6633 -1008 97 33 C
ATOM 466 O GLY A 63	16.677 -11.199 19.842 1.00 46.14 O

TABLE 3-continued

ANISOU 466 O GLY A 63	5412 6053 6066 -1042 97 168 O
ATOM 467 N TYR A 64	16.583 -9.428 18.506 1.00 47.27 N
ANISOU 467 N TYR A 64	5428 6290 6242 -870 49 -66 N
ATOM 468 CA TYR A 64	18.002 -9.087 18.731 1.00 46.70 C
ANISOU 468 CA TYR A 64	5458 6235 6049 -769 8 -29 C
ATOM 469 C TYR A 64	18.071 -7.933 19.686 1.00 45.48 C
ANISOU 469 C TYR A 64	5267 6231 5784 -763 96 -59 C
ATOM 470 O TYR A 64	17.051 -7.297 19.907 1.00 42.82 O
ANISOU 470 O TYR A 64	4820 5976 5472 -804 175 -133 O
ATOM 471 CB TYR A 64	18.703 -8.717 17.422 1.00 44.76 C
ANISOU 471 CB TYR A 64	5221 5945 5839 -630 -96 -103 C
ATOM 472 CG TYR A 64	18.781 -9.207 16.546 1.00 41.30 C
ANISOU 472 CG TYR A 64	4832 5374 5487 -613 -193 -98 C
ATOM 473 CD1 TYR A 64	19.788 -10.857 16.706 1.00 41.22 C
ANISOU 473 CD1 TYR A 64	4934 5275 5451 -582 -255 -21 C
ATOM 474 CD2 TYR A 64	17.786 -10.138 15.607 1.00 45.04 C
ANISOU 474 CD2 TYR A 64	5232 5804 6079 -626 -234 -186 C
ATOM 475 CE1 TYR A 64	19.826 -12.000 15.927 1.00 45.63 C
ANISOU 475 CE1 TYR A 64	5538 5693 6108 -557 -355 -41 C
ATOM 476 CE2 TYR A 64	17.608 -11.285 14.820 1.00 48.41 C
ANISOU 476 CE2 TYR A 64	5700 6099 6595 -611 -338 -212 C
ATOM 477 CZ TYR A 64	18.820 -12.205 14.997 1.00 44.62 C
ANISOU 477 CZ TYR A 64	5337 5521 6096 -577 -394 -143 C
ATOM 478 OH TYR A 64	18.806 -13.287 14.162 1.00 51.69 O
ANISOU 478 OH TYR A 64	6267 6276 7096 -545 -509 -202 O
ATOM 479 N LYS A 65	19.253 -7.662 20.236 1.00 43.57 N
ANISOU 479 N LYS A 65	5103 6024 5427 -705 72 -21 N
ATOM 480 CA LYS A 65	19.410 -6.561 21.182 1.00 47.04 C
ANISOU 480 CA LYS A 65	5512 6600 5761 -692 134 -76 C
ATOM 481 C LYS A 65	19.407 -5.217 20.490 1.00 45.63 C
ANISOU 481 C LYS A 65	5250 6434 5852 -609 113 -207 C
ATOM 482 O LYS A 65	19.111 -4.185 21.137 1.00 46.32 O
ANISOU 482 O LYS A 65	5276 6615 5707 -603 167 -299 O
ATOM 483 CB LYS A 65	20.662 -6.699 22.033 1.00 48.73 C
ANISOU 483 CB LYS A 65	5825 6853 5837 -659 96 -9 C
ATOM 484 CG LYS A 65	20.467 -7.697 23.156 1.00 62.37 C
ANISOU 484 CG LYS A 65	7629 8620 7448 -748 147 128 C
ATOM 485 CD LYS A 65	21.581 -8.712 23.185 1.00 64.02 C
ANISOU 485 CD LYS A 65	7965 8735 7626 -708 44 254 C
ATOM 486 CE LYS A 65	21.072 -9.958 23.895 1.00 72.21 C

TABLE 3-continued

ANISOU 486 CE LYS A 65	9081 9736 8620 -817 85 425 C
ATOM 487 NZ LYS A 65	21.402 -11.183 23.112 1.00 69.91 N
ANISOU 487 NZ LYS A 65	6862 9241 8458 -802 -17 508 N
ATOM 488 N GLY A 66	19.718 -5.211 19.186 1.00 40.53 N
ANISOU 488 N GLY A 66	4604 5695 5100 -540 32 -217 N
ATOM 489 CA GLY A 66	19.985 -3.947 18.496 1.00 35.60 C
ANISOU 489 CA GLY A 66	3928 50666 4532 -456 1 -294 C
ATOM 490 C GLY A 66	20.631 -4.208 17.139 1.00 37.45 C
ANISOU 490 C GLY A 66	4192 5224 4812 -380 -81 -259 C
ATOM 491 O GLY A 66	20.724 -5.361 16.711 1.00 35.04 O
ANISOU 491 O GLY A 66	3937 4868 4507 -383 -119 -212 O
ATOM 492 N TYR A 67	21.082 -3.145 16.462 1.00 32.81 N
ANISOU 492 N TYR A 67	3573 4629 4263 -311 -107 -282 N
ATOM 493 CA TYR A 67	21.595 -3.286 15.099 1.00 31.31 C
ANISOU 493 CA TYR A 67	3401 4403 4092 -233 -165 -246 C
ATOM 494 C TYR A 67	22.866 -4.135 15.076 1.00 32.53 C
ANISOU 494 C TYR A 67	3622 4554 4183 -206 -196 -186 C
ATOM 495 O TYR A 67	23.028 -5.041 14.230 1.00 29.85 O
ANISOU 495 O TYR A 67	3317 4189 3835 -159 -240 -172 O
ATOM 496 CB TYR A 67	21.865 -1.912 14.489 1.00 31.93 C
ANISOU 496 CB TYR A 67	3435 4476 4220 -180 -171 -247 C
ATOM 497 CG TYR A 67	22.476 -2.041 13.154 1.00 34.71 C
ANISOU 497 CG TYR A 67	3807 4828 4554 -103 -209 -189 C
ATOM 498 CD1 TYR A 67	21.714 -2.557 12.093 1.00 34.38 C
ANISOU 498 CD1 TYR A 67	3767 4787 4510 -56 -250 -204 C
ATOM 499 CD2 TYR A 67	23.859 -1.790 12.944 1.00 33.24 C
ANISOU 499 CD2 TYR A 67	3630 4661 4340 -76 -206 -126 C
ATOM 500 CE1 TYR A 67	22.271 -2.751 10.831 1.00 36.50 C
ANISOU 500 CE1 TYR A 67	4058 5088 4724 30 -284 -162 C
ATOM 501 CE2 TYR A 67	24.420 -1.962 11.665 1.00 33.19 C
ANISOU 501 CE2 TYR A 67	3631 4688 4291 1 -221 -71 C
ATOM 502 CZ TYR A 67	23.602 -2.415 10.605 1.00 31.81 C
ANISOU 502 CZ TYR A 67	3471 4527 4090 59 -258 -90 C
ATOM 503 OH TYR A 67	24.083 -2.694 9.382 1.00 33.21 O
ANISOU 503 OH TYR A 67	3662 4767 4188 150 -275 -54 O
ATOM 504 N SER A 68	23.816 -3.812 15.961 1.00 33.40 N
ANISOU 504 N SER A 68	3744 4693 4255 -220 -186 -168 N
ATOM 505 CA SER A 68	25.107 -4.557 15.900 1.00 31.87 C
ANISOU 505 CA SER A 68	3594 4502 4013 -174 -228 -119 C
ATOM 506 C SER A 68	24.901 -6.076 16.126 1.00 30.50 C

TABLE 3-continued

ANISOU 506 C SER A 68	3495 4283 3811 -184 -260 -89 C
ATOM 507 O SER A 68	25.576 -6.864 15.531 1.00 29.83 O
ANISOU 507 O SER A 68	3441 4171 3721 -117 -310 -74 O
ATOM 508 CB SER A 68	25.086 -3.991 16.913 1.00 34.59 C
ANISOU 508 CB SER A 68	3925 4890 4326 -190 -230 -119 C
ATOM 509 OG SER A 68	28.574 -2.710 16.439 1.00 38.20 O
ANISOU 509 OG SER A 68	4311 5358 4847 -175 -218 -134 O
ATOM 510 N ADP A 69	24.069 -6.455 17.107 1.00 32.20 N
ANISOU 510 N ASP A 69	3737 4490 4008 -269 -229 -77 N
ATOM 511 CA ASP A 69	23.701 -7.883 17.382 1.00 32.55 C
ANISOU 511 CA ASP A 69	3854 4460 4053 -308 -254 -23 C
ATOM 512 C ASP A 69	23.075 -8.574 16.143 1.00 32.67 C
ANISOU 512 C ASP A 69	3864 4395 4155 -282 -296 -61 C
ATOM 513 O ASP A 69	23.470 -9.655 15.688 1.00 36.11 O
ANISOU 513 O ASP A 69	4355 4749 4618 -236 -366 -50 O
ATOM 514 CB ASP A 69	22.656 -7.866 18.490 1.00 32.49 C
ANISOU 514 CB ASP A 69	3841 4484 4018 -423 -179 O C
ATOM 515 CG ASP A 69	22.352 -9.265 18.990 1.00 38.30 C
ANISOU 515 CG ASP A 69	4656 5140 4756 -490 -192 96 C
ATOM 516 OD1 ASP A 69	23.237 -10.180 18.949 1.00 37.71 O
ANISOU 516 OD1 ASP A 65	4654 4989 4575 -436 -272 157 O
ATOM 517 OD2 ASP A 69	21.201 -9.473 19.428 1.00 35.88 O
ANISOU 517 OD2 ASP A 69	4323 4840 4470 -598 -122 114 O
ATOM 518 N PHE A 70	22.047 -7.934 15.620 1.00 31.66 N
AMISOU 518 N PHE A 70	3664 4289 4076 -305 -264 -122 N
ATOM 519 CA PHE A 70	21.459 -8.300 14.296 1.00 33.63 C
ANISOU 519 CA PHE A 70	3890 4495 4392 -259 -320 -187 C
ATOM 520 C PHE A 70	22.500 -8.447 13.188 1.00 31.74 C
ANISOU 520 C PHE A 70	3675 4264 4119 -129 -382 -207 C
ATOM 521 O PHE A 70	22.519 -9.468 12.465 1.00 37.08 O
ANISOU 521 O PHE A 70	4387 4879 4824 -79 -455 -249 O
ATOM 522 CB PHE A 70	20.416 -7.252 13.916 1.00 35.08 C
ANISOU 522 CB PHE A 70	3983 4732 4615 -271 -287 -247 C
ATOM 523 CG PHE A 70	19.808 -7.481 12.561 1.00 36.66 C
ANISOU 523 CG PHE A 70	4154 4915 4862 -210 -358 -319 C
ATOM 524 CD1 PHE A 70	20.460 -7.053 11.431 1.00 38.73 C
ANISOU 524 CD1 PHE A 70	4424 5222 5071 -90 -399 -329 C
ATOM 525 CD2 PHE A 70	18.549 -8.107 12.447 1.00 42.13 C
ANISOU 525 CD2 PHE A 70	4801 5561 5647 -277 -383 -376 C
ATOM 526 CE1 PHE A 70	19.903 -7.287 10.161 1.00 36.96 C

TABLE 3-continued

ANISOU 526 CE1 PHE A 70	4181 5007 4855 -17 -476 -402 C
ATOM 527 CE2 PHE A 70	17.976 -8.340 11.201 1.00 46.07 C
ANISOU 527 CE2 PHE A 70	5268 6053 6184 -213 -472 -465 C
ATOM 528 CZ PHE A 70	18.659 -7.900 10.055 1.00 43.17 C
ANISOU 528 CZ PHE A 70	4925 5744 5733 -73 -522 -480 C
ATOM 529 N ARG A 71	23.383 -7.424 13.024 1.00 33.14 N
ANISOU 529 N ARG A 71	3827 4523 4243 -72 -352 -185 N
ATOM 530 CA ARG A 71	24.316 -7.446 11.903 1.00 30.96 C
ANISOU 530 CA ARG A 71	3550 4291 3922 46 -383 -196 C
ATOM 531 C ARG A 71	25.306 -8.640 12.082 1.00 30.94 C
ANISOU 531 C ARG A 71	3605 4248 3904 100 -435 -192 C
ATOM 532 O ARG A 71	25.664 -9.306 11.142 1.00 31.69 O
ANISOU 532 O ARG A 71	3714 4344 3984 199 -485 -245 O
ATOM 533 CB ARG A 71	25.076 -6.138 11.831 1.00 31.36 C
ANISOU 533 CB ARG A 71	3550 4422 3942 65 -330 -149 C
ATOM 534 CG ARG A 71	25.762 -5.929 10.464 1.00 37.50 C
ANISOU 534 CG ARG A 71	4305 5280 4664 175 -332 -143 C
ATOM 535 CD ARG A 71	26.529 -4.605 10.537 1.00 39.36 C
ANISOU 535 CD ARG A 71	4482 5571 4902 158 -270 -69 C
ATOM 536 NE ARG A 71	27.790 -4.838 11.252 1.00 33.84 N
ANISOU 536 NE ARG A 71	3769 4889 4198 155 -263 -51 N
ATOM 537 CZ ARG A 71	28.960 -4.212 11.034 1.00 40.34 C
ANISOU 537 CZ ARG A 71	4527 5782 5018 174 -223 -2 C
ATOM 538 NH1 ARG A 71	29.062 -3.248 10.125 1.00 40.55 N
ANISOU 538 NH1 ARG A 71	4506 5860 5041 184 -173 60 N
ATOM 539 NH2 ARG A 71	30.049 -4.524 11.779 1.00 38.05 N
ANISOU 539 NH2 ARG A 71	4214 5511 4737 175 -238 -6 N
ATOM 540 N MET A 72	25.740 -8.883 13.321 1.00 30.52 N
ANISOU 540 N MET A 72	3585 4163 3846 46 -429 -137 N
ATOM 541 CA MET A 72	26.597 -10.058 13.611 1.00 31.07 C
ANISOU 541 CA MET A 72	3716 4171 3919 103 -496 -122 C
ATOM 542 C MET A 72	25.877 -11.384 13.207 1.00 37.86 C
ANISOU 542 C MET A 72	4636 4897 4852 106 -570 -166 C
ATOM 543 O MET A 72	26.468 -12.243 12.501 1.00 35.91 O
ANISOU 543 O MET A 72	4414 4608 4623 219 -644 -226 O
ATOM 544 CB MET A 72	26.866 -10.086 15.136 1.00 30.13 C
ANISOU 544 CB MET A 72	3635 4039 3773 29 -487 -39 C
ATOM 545 CB MET A 72	27.719 -11.262 15.653 1.00 54.11 C
ANISOU 545 CB MET A 72	4213 4467 4279 97 -572 5 C
ATOM 546 SD MET A 72	29.522 -11.016 15.312 1.00 37.82 S

TABLE 3-continued

ANISOU 546 SD MET A 72	4622 5036 4713 234 -608 -26 S
ATOM 547 CE MET A 72	29 591 -12.134 13.917 1.00 36.99 C
ANISOU 547 CE MET A 72	4533 4857 4664 365 -675 -121 C
ATOM 548 N ALA A 73	24.621 -11.570 13.671 1.00 34.56 N
ANISOU 548 N ALA A 73	4231 4410 4490 -17 -551 -146 N
ATOM 549 CA ALA A 73	23.880 -22.799 13.245 1.00 37.46 C
ANISOU 549 CA ALA A 73	4638 4632 4962 -36 -627 -194 C
ATOM 550 C ALA A 73	23.775 -12.849 11.743 1.00 38.12 C
ANISOU 550 C ALA A 73	4687 4745 5053 75 -682 -329 C
ATOM 551 O ALA A 73	23.839 -13.947 11.110 1.00 40.11 O
ANISOU 551 O ALA A 73	4978 4891 5370 143 -783 -415 O
ATOM 552 CB ALA A 73	22.483 -12.825 13.835 1.00 39.86 -161 C
ANISOU 552 CB ALA A 73	4921 4888 5335 -197 -580 -161 C
ATOM 553 N LEU A 74	23.552 -11.684 11.154 1.00 37.81 N
ANISOU 553 N LEU A 74	4577 4841 4947 97 -627 -354 N
ATOM 554 CA LEU A 74	23.447 -11.660 9 707 1 00 36.77 C
ANISOU 554 CA LEU A 74	4419 4770 4782 211 -677 -466 C
ATOM 555 C LEU A 74	24.701 -12.260 9.058 1.00 38.94 C
ANISOU 555 C LEU A 74	4722 5073 5001 366 -725 -520 C
ATOM 556 O LEU A 74	24.606 -13.148 8.151 1.00 35.78 O
ANISOU 556 O LEU A 74	4341 4633 4619 460 -818 -650 O
ATOM 557 CB LEU A 74	23.183 -10.232 9.208 1.00 36.57 C
ANISOU 557 CB LEU A 74	4326 4888 4682 223 -611 -444 C
ATOM 558 CG LEU A 74	23.199 -10.096 7.682 1.00 36.01 C
ANISOU 558 CG LEU A 74	4236 4919 4527 357 -655 -529 C
ATOM 559 CD1 LEU A 74	22.052 -10.938 7.032 1.00 40.19 C
ANISOU 559 CD1 LEU A 74	4767 5378 5126 361 -764 -664 C
ATOM 560 CD2 LEU A 74	23.028 -8.655 7.261 1.00 34.53 C
ANISOU 560 CD2 LEU A 74	3997 4853 4269 367 -591 -463 C
ATOM 561 N ALA A 75	25.884 -11.789 9.504 1.00 36.22 N
ANISOU 561 N ALA A 75	4366 4804 4593 400 -567 -443 N
ATOM 562 CA ALA A 75	27.147 -12.187 8.845 1.00 39.53 C
ANISOU 562 CA ALA A 75	4776 5294 4951 557 -690 -498 C
ATOM 563 C ALA A 75	27.362 -13.696 9.059 1.00 42.03 C
ANISOU 563 C ALA A 75	5163 5447 5360 610 -803 -567 C
ATOM 564 O ALA A 75	27.808 -14.408 8.170 1.00 41.98 O
ANISOU 564 O ALA A 75	5160 5451 5340 755 -868 -695 O
ATOM 565 CB ALA A 75	28.359 -11.332 9.364 1.00 35.99 C
ANISOU 565 CB ALA A 75	4274 4958 4443 564 -608 -401 C
ATOM 566 N VAL A 76	27.018 -14.132 10.251 1.00 43.81 N

TABLE 3-continued

ANISOU 566 N VAL A 76	5448 5518 5679 496 -827 -482 N
ATOM 567 CA VAL A 76	27.175 -15.596 10.573 1.00 43.37 C
ANISOU 567 CA VAL A 76	5473 5268 5736 528 -941 -509 C
ATOM 568 C VAL A 76	26.211 -16.402 9.705 1.00 46.98 C
ANISOU 568 C VAL A 76	5952 5610 6290 5.36 -1031 -647 C
ATOM 569 O VAL A 76	26.621 -17.443 9.162 1.00 52.07 O
ANISOU 569 O VAL A 76	6632 6155 6998 661 -1142 -770 O
ATOM 570 CB VAL A 76	26.923 -15.875 12.058 1.00 40.62 C
ANISOU 570 CB VAL A 76	5191 4793 5446 386 -935 -349 C
ATOM 571 CG1 VAL A 76	26.894 -17.391 12.355 1.00 43.16 C
ANISOU 571 CG1 VAL A 76	5613 4870 5916 399 -1063 -348 C
ATOM 572 CG2 VAL A 76	28.011 -15.209 12.889 1.00 44.04 C
ANISOU 572 CG2 VAL A 76	5605 5341 5786 408 -883 -248 C
ATOM 573 N ASP A 77	24.978 -15.881 9.507 1.00 44.58 N
ANISOU 573 N ASP A 77	5612 5330 5997 421 -994 -652 N
ATOM 574 CA ASP A 77	23.928 -16.580 8.751 1.00 48.86 C
ANISOU 574 CA ASP A 77	6154 5765 6644 405 -1089 -791 C
ATOM 575 C ASP A 77	24.370 -16.673 7.266 1.00 54.24 C
ANISOU 575 C ASP A 77	6809 6565 7233 604 -1151 -982 C
ATOM 576 O ASP A 77	24.283 -17.762 6.631 1.00 56.34 O
ANISOU 576 O ASP A 77	7107 6710 7568 688 -1283 -1149 O
ATOM 577 CB ASP A 77	22.572 -15.870 8.919 1.00 52.98 C
ANISOU 577 CB ASP A 77	6617 6322 7192 249 -1030 -753 C
ATOM 578 CG ASP A 77	21.363 -16.768 8.624 1.00 66.58 C
ANISOU 578 CG ASP A 77	6331 7877 9088 164 -1132 -857 C
ATOM 579 OD1 ASP A 77	21.413 -18.010 8.900 1.00 73.09 O
ANISOU 579 OD1 ASP A 77	9220 8484 10067 139 -1226 -876 O
ATOM 580 OD2 ASP A 77	20.334 -16.206 8.125 1.00 59.17 O
ANISOU 580 OD2 ASP A 77	7315 7018 8150 119 -1126 -918 O
ATOM 581 N LEU A 78	24.908 -15.581 6.725 1.00 47.73 N
ANISOU 581 N LEU A 78	5931 5975 6231 684 -1060 -961 N
ATOM 582 CA LEU A 78	25.437 -15.582 5.350 1.00 52.40 C
ANISOU 582 CA LEU A 78	6494 6728 6686 877 -1088 -1112 C
ATOM 583 C LEU A 78	26.602 -16.552 5.173 1.00 53.92 C
ANISOU 583 C LEU A 78	6714 6883 6890 1037 -1150 -1212 C
ATOM 584 O LEU A 78	26.777 -17.148 4.117 1.00 60.12 O
ANISOU 584 O LEU A 78	7497 7714 7630 1197 -1229 -1405 O
ATOM 585 CB LEU A 78	25.857 -14.167 4.954 1.00 48.67 C
ANISOU 585 CB LEU A 78	5958 6503 6031 905 -955 -1012 C
ATOM 586 CG LEU A 78	24.695 -13.198 4.630 1.00 49.90 C

TABLE 3-continued

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ANISOU 586 CG LEU A 78	6081 6729 6150 821 -926 -971 C
ATOM 587 CD1 LEU A 78	25.210 -11.803 4.357 1.00 49.89 C
ANISOU 587 CD1 LEU A 78	6030 6927 6000 838 -799 -840 C
ATOM 588 CD2 LEU A 78	23.885 -13.683 3.444 1.00 52.17 C
ANISOU 588 CD2 LEU A 78	6370 7044 6409 908 -1042 -1154 C
ATOM 589 N SER A 79	27.428 -16.635 6.208 1.00 57.86 N
ANISOU 589 N SER A 79	7230 7319 7435 1008 -1113 -1087 N
ATOM 590 CA SER A 79	28.665 -17.407 6.239 1.00 61.23 C
ANISOU 590 CA SER A 79	7667 7719 7880 1163 -1163 -1152 C
ATOM 591 C SER A 79	28.609 -18.838 5.708 1.00 72.71 C
ANISOU 591 C SER A 79	9175 8997 9456 1289 -1328 -1358 C
ATOM 592 O SER A 79	29.630 -16.379 5.231 1.00 73.62 O
ANISOU 592 O SER A 79	9270 9160 9543 1484 -1370 -1468 O
ATOM 593 CB SER A 79	29.121 -17.511 7.676 1.00 59.22 C
ANISOU 593 CB SER A 79	7450 7339 7712 1070 -1152 -978 C
ATOM 594 OG SER A 79	30.421 -17.003 7.752 1.00 62.93 O
ANISOU 594 OG SER A 79	7850 7977 8082 1166 -1081 -939 O
ATOM 595 N GLN A 80	27.448 -19.470 5.834 1.00 74.00 N
ANISOU 595 N GLN A 80	9397 8947 9773 1177 -1425 -1396 N
ATOM 596 CA GLN A 80	27.291 -20.837 5.353 1.00 92.13 C
ANISOU 596 CA GLN A 80	11747 11033 12227 1275 -1601 -1602 C
ATOM 597 C GLN A 80	27.085 -20.893 3.831 1.00 98.84 C
ANISOU 597 C GLN A 80	12552 12040 12963 1436 -1655 -1863 C
ATOM 598 O GLN A 80	26.665 -21.926 3.306 1.00 103.30 O
ANISOU 598 O GLN A 80	13153 12436 13662 1499 -1815 -2073 O
ATOM 599 CB GLN A 80	26.148 -21.539 6.083 1.00 96.22 C
ANISOU 599 CB GLN A 80	12332 11247 12980 1077 -1686 -1537 C
ATOM 600 CG GLN A 80	25.854 -20.957 7.457 1.00 99.68 C
ANISOU 600 CG GLN A 80	12788 11647 13437 862 -1572 -1251 C
ATOM 601 CD GLN A 80	24.365 -20.796 7.711 1.00 96.67 C
ANISOU 601 CD GLN A 80	12393 11184 13153 643 -1556 -1195 C
ATOM 602 OE1 GLN A 80	23.954 -20.405 8.799 1.00 105.29 O
ANISOU 602 OE1 GLN A 80	13494 12247 14265 463 -1463 -987 O
ATOM 603 NE2 GLN A 80	23.553 -21.083 6.705 1.00 93.40 N
ANISOU 603 NE2 GLN A 80	11944 10751 12791 664 -1646 -1394 N
ATOM 604 N THR A 81	27.375 -19.786 3.136 1.00 98.15 N
ANISOU 604 N THR A 81	12391 12273 12630 1501 -1527 -1846 N
ATOM 605 CA THR A 81	27.430 -19.760 1.661 1.00 99.77 C
ANISOU 605 CA THR A 81	12555 12696 12656 1690 -1556 -2072 C
ATOM 606 C THR A 81	28.474 -18.762 1.150 1.00 96.99 C

TABLE 3-continued

ANISOU 606 C THR A 81	12124 12686 12042 1805 -1390 -2003 C
ATOM 607 O THR A 81	28.192 -17.932 0.280 1.00 88.18 O
ANISOU 607 O THR A 81	10968 11813 10722 1836 -1320 -2002 O
ATOM 608 CB THR A 81	26.060 -19.433 1.021 1.00101.17 C
ANISOU 608 CB THR A 81	12727 12909 12805 1611 -1606 -2137 C
ATOM 609 OG1 THR A 81	25.428 -18.377 1.762 1.00 96.54 O
ANISOU 609 OG1 THR A 81	12122 12340 12220 1404 -1492 -1891 O
ATOM 610 CG2 THR A 81	25.149 -20.682 0.964 1.00 96.71 C
ANISOU 610 CG2 THR A 81	12211 12051 12484 1575 -1810 -2832 C
ATOM 611 N GLY A 91	47.291 -17.018 -6.584 1.00 6976 N
ANISOU 611 N GLY A 91	9436 9270 7801 526 -543 -1048 N
ATOM 612 CA GLY A 91	46.882 -18.155 -5.748 1.00 71 02 C
ANISOU 612 CA GLY A 91	9610 9325 8050 466 -637 -1088 C
ATOM 613 C GLY A 91	47.505 -18.348 -4.355 1.00 71.89 C
ANISOU 613 C GLY A 91	9695 9331 8289 445 -626 -1059 C
ATOM 614 O GLY A 91	46.772 -18.620 -3.387 1.00 68.11 O
ANISOU 614 O GLY A 91	9202 8797 7878 373 -686 -1021 O
ATOM 615 N ASP A 92	48.837 -18.235 -4.241 1.00 65.60 N
ANISOU 615 N ASP A 92	8891 8509 7524 505 -552 -1079 N
ATOM 616 CA ASP A 92	49.583 -18.449 -2.975 1.00 63.49 C
ANISOU 616 CA ASP A 92	8603 8145 7376 496 -542 -1061 C
ATOM 617 C ASP A 92	49.182 -17.457 -1.815 1.00 63.40 C
ANISOU 617 C ASP A 92	8533 8135 7422 446 -518 -941 C
ATOM 618 O ASP A 92	48.943 -16.273 -2.030 1.00 55.84 O
ANISOU 618 O ASP A 92	7540 7258 6420 452 -459 -868 O
ATOM 619 CB ASP A 92	51.073 -18.510 -3.343 1.00 64.67 C
ANISOU 619 CB ASP A 92	8752 8289 7530 579 -468 -1119 C
ATOM 620 CG ASP A 92	52.023 -18.042 -2.268 1.00 68.95 C
ANISOU 620 CG ASP A 92	9246 8785 8166 590 -412 -1071 C
ATOM 621 OD1 ASP A 92	52.810 -18.869 -1.766 1.00 72.78 O
ANISOU 621 OD1 ASP A 92	9744 9184 8726 613 -434 -1127 O
ATOM 622 OD2 ASP A 92	52.108 -16.834 -2.007 1.00 57.33 O
ANISOU 622 OD2 ASP A 92	7725 7365 6691 587 -341 -984 O
ATOM 623 N ILE A 93	49.048 -17.968 -0.601 1.00 59.05 N
ANISOU 623 N ILE A 93	7979 7499 6966 395 -557 -922 N
ATOM 624 CA ILE A 93	48.528 -17.150 0.500 1.00 59.91 C
ANISOU 624 CA ILE A 93	8038 7602 7122 343 -555 -818 C
ATOM 625 C ILE A 93	49.311 -15.862 0.686 1.00 55.48 C
ANISOU 625 C ILE A 93	7427 7085 6566 384 -451 -755 C
ATOM 626 O ILE A 93	48 728 -14.847 0.967 1.00 56 94 O

TABLE 3-continued

ANISOU 626 O ILE A 93	7575 7321 6740 361 -423 -672 O
ATOM 627 CB ILE A 93	48.361 -17.954 1.807 1.00 63.95 C
ANISOU 627 CB ILE A 93	8559 8007 7732 284 -621 -809 C
ATOM 628 CG1 ILE A 93	47.427 -17.224 2.783 1.00 69.15 C
ANISOU 628 CG1 ILE A 93	9173 8684 8417 217 -624 -708 C
ATOM 629 CG2 ILE A 93	49.715 -18.244 2.448 1.00 70.74 C
ANISOU 629 CG2 ILE A 93	9420 8791 8667 327 -593 -834 C
ATOM 630 CD1 ILE A 93	46.038 -17.000 2.215 1.00 76.48 C
ANISOU 630 CD1 ILE A 93	10093 9687 9279 173 -660 -686 C
ATOM 631 N CYS A 94	50.620 -15.870 0.466 1.00 55.79 N
ANISOU 631 N CYS A 94	7465 7112 6620 447 -393 -796 N
ATOM 632 CA CYS A 94	51.360 -14.631 0.482 1.00 53.36 C
ANISOU 632 CA CYS A 94	7112 6854 6310 483 -292 -740 C
ATOM 633 C CYS A 94	50.890 -13.659 -0.550 1.00 55.68 C
ANISOU 633 C CYS A 94	7398 7253 6504 500 -242 -702 C
ATOM 634 O CYS A 94	50.787 -12.464 -0.266 1.00 48.81 O
ANISOU 634 O CYS A 94	6489 6421 5634 493 -187 -617 O
ATOM 635 CB CYS A 94	52.842 -14.848 0.208 1.00 55.56 C
ANISOU 635 CB CYS A 94	7385 7114 6610 550 -235 -801 C
ATOM 636 SG CYS A 94	53.702 -15.632 1.584 1.00 68.24 S
ANISOU 636 SG CYS A 94	8963 8600 9346 547 -271 -826 S
ATOM 637 N ASP A 95	50.692 -14.153 -1.770 1.00 53.56 N
ANISOU 637 N ASP A 95	7171 7029 6150 527 -260 -766 N
ATOM 638 CA ASP A 95	50.453 -13.276 -2.899 1.00 52.88 C
ANISOU 638 CA ASP A 95	7086 7047 5958 557 -207 -737 C
ATOM 639 C ASP A 95	49.063 -12.689 -2.806 1.00 48.80 C
ANISOU 639 C ASP A 95	6559 6574 5409 510 -247 -663 C
ATOM 640 O ASP A 95	48.838 -11.533 -3.151 1.00 47.88 O
ANISOU 640 O ASP A 95	6423 6527 5244 522 -193 -591 O
ATOM 641 CB ASP A 95	50 664 -14.016 -4.233 1.00 59.07 C
ANISOU 641 CB ASP A 95	7921 7870 6552 603 -217 -833 C
ATOM 642 CG ASP A 95	52.150 -14.410 -4.466 1.00 63.52 C
ANISOU 642 CG ASP A 95	8486 8411 7238 665 -156 -905 C
ATOM 643 OD1 ASP A 95	52.407 -15.613 -4.688 1.00 65.12 O
ANISOU 643 OD1 ASP A 95	8726 8566 7450 683 -207 -1004 O
ATOM 644 OD2 ASP A 95	53.052 -13.540 -4.370 1.00 62.19 O
ANISOU 644 OD2 ASP A 95	8278 8266 7085 693 -61 -864 O
ATOM 645 N VAL A 96	48.109 -13.481 -2.353 1.00 45.40 N
ANISOU 645 N VAL A 96	6140 6104 5006 455 -342 -680 N
ATOM 646 CA VAL A 96	46.755 -12.953 -2.148 1.00 44.62 C

TABLE 3-continued

ANISOU 646 CA VAL A 96	6020 6047 4888 407 -382 -611 C
ATOM 647 C VAL A 95	46.694 -11.948 -0.985 1.00 42.69 C
ANISOU 647 C VAL A 95	5721 5786 4712 383 -341 -515 C
ATOM 648 O VAL A 96	46.031 -10.888 -1.088 1.00 42.06 O
ANISOU 648 O VAL A 96	5615 5769 4598 381 -318 -440 O
ATOM 649 CB VAL A 96	45.721 -14.072 -1.928 1.00 48.61 C
ANISOU 649 CB VAL A 96	6544 6516 5409 346 -493 -653 C
ATOM 650 CG1 VAL A 96	44.317 -13.457 -1 969 1.00 48.28 C
ANISOU 650 CG1 VAL A 96	6472 6541 5330 307 -529 -589 C
ATOM 651 CG2 VAL A 96	45.870 -15.189 -3.004 1.00 46.43 C
ANISOU 651 OG2 VAL A 96	6327 6239 5075 370 -540 -763 C
ATOM 652 N SER A 97	47.407 -12.211 0.106 1.00 40.76 N
ANISOU 652 N SER A 97	5463 5461 4563 369 -331 -515 N
ATOM 653 CA SER A 97	47.282 -11.318 1.248 1.00 41.24 C
ANISOU 653 CA SER A 97	5477 5506 4686 342 -300 -430 C
ATOM 654 C SER A 97	47.895 -9.985 0.850 1.00 40.75 C
ANISOU 654 C SER A 97	5393 5497 4594 390 -202 -377 C
ATOM 655 O SER A 97	47.331 -8.914 1.128 1.00 40.30 O
ANISOU 655 O SER A 97	5305 5475 4532 381 -176 -296 O
ATOM 656 CB SER A 97	47.994 -11.887 2.486 1.00 42.57 C
ANISOU 656 CB SER A 97	5640 5578 4957 321 -311 -444 C
ATOM 657 OG SER A 97	47.477 -13.186 2.822 1.00 45.48 O
ANISOU 657 OG SER A 97	6039 5889 5354 275 -402 -493 O
ATOM 658 N ALA A 98	49.040 -10.045 0.147 1.00 42.05 N
ANISOU 658 N ALA A 98	5573 5668 4735 443 -146 -421 N
ATOM 659 CA ALA A 98	49.715 -8.825 -0.367 1.00 41.79 C
ANISOU 659 CA ALA A 98	5524 5688 4667 486 -46 -374 C
ATOM 660 C ALA A 98	48.846 -8.097 -1.402 1.00 45.61 C
ANISOU 660 C ALA A 98	6021 6261 5049 500 -39 -330 C
ATOM 661 O ALA A 98	48.696 -6.833 -1.349 1.00 41.70 O
ANISOU 661 O ALA A 98	5503 5798 4543 507 15 -247 O
ATOM 662 CB ALA A 98	51.050 -9.185 -1.026 1.00 44.11 C
ANISOU 662 CB ALA A 98	5831 5979 4949 536 10 -439 C
ATOM 663 N GLN A 99	48.231 -8.864 -2.309 1.00 45.74 N
ANISOU 663 N GLN A 99	6075 6314 4992 505 -97 -385 N
ATOM 664 CA GLN A 99	47.375 -8.205 -3.304 1.00 43.56 C
ANISOU 664 CA GLN A 99	5813 6126 4613 521 -100 -345 C
ATOM 665 C GLN A 99	46.189 -7.447 -2.698 1.00 41.45 C
ANISOU 665 C GLN A 99	5512 5875 4362 488 -131 -262 C
ATOM 666 O GLN A 99	45.925 -6.312 -3.139 1.00 35.07 O

TABLE 3-continued

ANISOU 666 O GLN A 99	4699 5123 3504 513 -89 -192 O
ATOM 667 CB GLN A 99	46.939 -9.127 -4.443 1.00 47.48 C
ANISOU 667 CB GLN A 99	6356 6666 5019 534 -159 -421 C
ATOM 668 CG GLN A 99	46.252 -8.392 -5.635 1.00 46.15 C
ANISOU 668 CG GLN A 99	6207 6598 4728 565 -153 -382 C
ATOM 669 CD GLN A 99	47.218 -7.514 -6.431 1.00 49.52 C
ANISOU 669 CD GLN A 99	6649 7072 5094 620 -47 -352 C
ATOM 670 OE1 GLN A 99	48.256 -7.978 -6.909 1.00 47.86 O
ANISOU 670 OE1 GLN A 99	6461 6858 4866 650 -2 -414 O
ATOM 671 NE2 GLN A 99	45.880 -6.228 -6.584 1.00 49.20 N
ANISOU 671 NE2 GLN A 99	6597 7076 5020 633 -4 -255 N
ATOM 672 N SER A 100	45.508 -8.008 -1.677 1.00 39.71 N
ANISOU 672 N SER A 100	5269 5606 4213 433 -199 -267 N
ATOM 673 CA SER A 100	44.415 -7.300 -1.069 1.00 42.09 C
ANISOU 673 CA SER A 100	5533 5928 4532 404 -222 -194 C
ATOM 674 C SER A 100	44.900 -5.956 -0.549 1.00 39.63 C
ANISOU 674 C SER A 100	5192 5610 4255 425 -139 -113 C
ATOM 675 O SER A 100	44.193 -4.956 -0.654 1.00 38.14 O
ANISOU 675 O SER A 100	4986 5467 4038 436 -128 -44 O
ATOM 676 CB SER A 100	43.846 -8.060 0.125 1.00 47.00 O
ANISOU 676 CB SER A 100	6131 6491 5236 340 -268 -207 C
ATOM 677 OG SER A 100	43.222 -9.208 -0.338 2.00 51.97 O
ANISOU 677 OG SER A 100	6784 7126 5835 312 -371 -272 O
ATOM 678 N ALA A 101	46.030 -5.994 0.178 1.00 41.18 N
ANISOU 678 N ALA A 101	5381 5741 4525 423 -90 -124 N
ATOM 679 CA ALA A 101	46.554 -4.803 0.817 1.00 37.26 C
ANISOU 679 CA ALA A 101	4856 5227 4076 434 -16 -56 C
ATOM 680 C ALA A 101	46.937 -3.793 -9.259 1.00 40.05 C
ANISOU 680 C ALA A 101	5226 5636 4355 484 57 -15 C
ATOM 681 O ALA A 101	46.683 -2 594 -0.089 1.00 41.58 O
ANISOU 681 O ALA A 101	5404 5845 4551 494 95 61 O
ATOM 682 CB ALA A 101	47.735 -5.169 1.739 1.00 42.54 C
ANISOU 682 CB ALA A 101	5512 5818 4835 422 13 -86 C
ATOM 683 N VAL A 102	47.497 -4.260 -1.390 1.00 36.65 N
ANISOU 683 N VAL A 102	4831 5236 3855 516 74 -65 N
ATOM 684 CA VAL A 102	47.756 -3.349 -2.530 1.00 39.26 C
ANISOU 684 CA VAL A 102	5186 5633 4099 561 140 -24 C
ATOM 685 C VAL A 102	46.522 -2.651 -2.999 1.00 40.19 C
ANISOU 685 C VAL A 102	5310 5811 4151 571 108 38 C
ATOM 686 O VAL A 102	46.535 -1.411 -3.156 1.00 37.37 O

TABLE 3-continued

ANISOU 686 O VAL A 102	4952 5472 3776 592 164 117 O
ATOM 687 CB VAL A 102	48.460 -4.031 -3.718 1.00 39.32 C
ANISOU 687 CB VAL A 102	5233 5677 4029 595 162 -93 C
ATOM 688 CG1 VAL A 102	48 489 -3.120 -4.946 1.00 37.65 C
ANISOU 688 CG1 VAL A 102	5053 5542 3710 636 219 -42 C
ATOM 689 CG2 VAL A 102	49.899 -4.350 -3.290 1.00 37.59 C
ANISOU 689 CG2 VAL A 102	4999 5404 3879 598 221 -137 C
ATOM 690 N ASP A 103	45.440 -3.421 -3.189 1.00 44.61 N
ANISOU 690 N ASP A 103	5874 6397 4677 554 16 3 N
ATOM 691 CA ASP A 103	44.176 -2.892 -3.785 1.00 44.64 C
ANISOU 691 CA ASP A 103	5882 6472 4608 568 -29 51 C
ATOM 692 C ASP A 103	43.638 -1.882 -2.770 1.00 39.39 C
ANISOU 692 C ASP A 103	5173 5783 4010 555 -19 131 C
ATOM 693 O ASP A 103	43.208 -0.820 -3.130 1.00 39.33 O
ANISOU 693 O ASP A 103	5167 5813 3963 586 1 202 O
ATOM 694 CB ASP A 103	43.094 -3.991 -3.950 1.00 44.25 C
ANISOU 694 CB ASP A 103	5832 6447 4534 539 -138 -8 C
ATOM 695 CG ASP A 103	43.476 -5.065 -4.952 1.00 48.96 C
ANISOU 695 CG ASP A 103	6475 7065 5061 551 -152 -97 C
ATOM 696 OD1 ASP A 103	44.344 -4.819 -5.777 1.00 44.27 O
ANISOU 696 OD1 ASP A 103	5916 6495 4409 593 -98 -103 O
ATOM 697 OD2 ASP A 103	42.936 -6.175 -4.853 1.00 53.52 O
ANISOU 697 OD2 ASP A 103	7056 7632 5649 516 -243 -162 O
ATOM 698 N SER A 104	43.666 -2.244 -1.497 1.00 40.29 N
ANISOU 698 N SER A 104	5250 5833 4224 512 -36 117 N
ATOM 699 CA SER A 104	43.239 -1.291 -0.478 1.00 40.35 C
ANISOU 699 CA SER A 104	5218 5818 4297 501 -20 185 C
ATOM 700 C SER A 104	43.967 0.042 -0.405 1.00 38.02 C
ANISOU 700 C SER A 104	4924 5503 4017 533 70 254 C
ATOM 701 O SER A 104	43.343 1.080 -0.118 1.00 41.45 O
ANISOU 701 O SER A 104	5342 5948 4461 547 78 322 O
ATOM 702 CB SER A 104	43.245 -1.955 0.885 1.00 39.14 C
ANISOU 702 CB SER A 104	5030 5599 4242 448 -49 156 C
ATOM 703 OG SER A 104	42.144 -2.608 0.848 1.00 52.58 O
ANISOU 703 OG SER A 104	6722 7331 5926 417 -137 123 O
ATOM 704 N LEU A 105	45.285 -0.002 -0.530 1.00 37.01 N
ANISOU 704 N LEU A 105	4814 5343 3906 539 137 233 N
ATOM 705 CA LEU A 105	46.099 1.215 -0.571 1.00 34.88 C
ANISOU 705 CA LEU A 105	4549 5054 3548 562 228 294 C
ATOM 706 C LEU A 105	45.706 2.032 -1.756 1.00 38.10 C

TABLE 3-continued

ANISOU 706 C LEU A 105	4992 5526 3960 606 247 351 C
ATOM 707 O LEU A 105	45.492 3.224 -1.659 1.00 37.80 O
ANISOU 707 O LEU A 105	4953 5483 3926 625 281 427 O
ATOM 708 CB LEU A 105	47.582 0.854 -0.619 1.00 37.21 C
ANISOU 708 CB LEU A 105	4852 5315 3972 559 291 250 C
ATOM 709 CG LEU A 105	48.135 0.349 0.735 1.00 34.89 C
ANISOU 709 CG LEU A 105	4521 4945 3789 520 283 213 C
ATOM 710 CD1 LEU A 105	49.463 -0.347 0.461 1.00 35.96 C
ANISOU 710 CD1 LEU A 105	4664 5062 3938 525 322 149 C
ATOM 711 CD2 LEU A 105	48.370 1.559 1.635 1.00 34.19 C
ANISOU 711 CD2 LEU A 105	4406 4813 3771 513 333 278 C
ATOM 712 N GLN A 106	45.574 1.400 -2.905 1.00 39.15 N
ANISOU 712 N GLN A 106	5160 5717 3998 626 223 314 N
ATOM 713 CA GLN A 106	45.282 2.194 -4.087 1.00 41.21 C
ANISOU 713 CA GLN A 106	5461 6041 4156 671 244 372 C
ATOM 714 C GLN A 106	43.872 2.765 -4.007 1.00 40.26 C
ANISOU 714 C GLN A 106	5327 5953 4017 686 182 426 C
ATOM 715 O GLN A 106	43.668 3.946 -4.292 1.00 42.89 O
ANISOU 715 O GLN A 106	5675 6297 4324 719 215 508 O
ATOM 716 CB GLN A 106	45.491 1.365 -5.333 1.00 40.37 C
ANISOU 716 CC GLN A 106	5398 5995 3947 690 232 314 C
ATOM 717 CG GLN A 106	46.943 0.896 -5.474 1.00 44.75 C
ANISOU 717 CG GLN A 106	5962 6524 4518 685 304 262 C
ATOM 718 CD GLN A 106	47.120 -0.049 -6.647 1.00 50.84 C
ANISOU 718 CD GLN A 106	6774 7352 5189 705 287 190 C
ATOM 719 OE1 GLN A 106	46.214 -0.837 -7.027 1.00 50.30 O
ANISOU 719 OE1 GLN A 106	6718 7323 5072 705 199 144 O
ATOM 720 NE2 GLN A 106	48.278 0.037 -7.244 1.00 49.57 N
ANISOU 720 NE2 GLN A 106	6634 7202 4997 723 372 177 N
ATOM 721 N ASP A 107	42.913 1.952 -3.592 1.00 41.68 N
ANISOU 721 N ASP A 107	5477 6145 4214 661 94 382 N
ATOM 722 CA ASP A 107	41.573 2.488 -3.335 1.00 43.13 C
ANISOU 722 CA ASP A 107	5631 6358 4397 672 36 430 C
ATOM 723 C ASP A 107	41.530 3.630 -2.351 1.00 41.97 C
ANISOU 723 C ASP A 107	5456 6162 4330 676 76 498 C
ATOM 724 O ASP A 107	40.854 4.610 -2.620 1.00 40.76 O
ANISOU 724 O ASP A 107	5304 6035 4147 715 72 566 O
ATOM 725 CB ASP A 107	40.617 1.386 -2.900 1.00 43.00 C
ANISOU 725 CB ASP A 107	5578 6359 4402 632 -59 369 C
ATOM 726 CG ASP A 107	40.334 0.416 -4.019 1.00 49.56 C

TABLE 3-continued

ANISOU 726 CG ASP A 107	6440 7250 5140 637 -117 309 C
ATOM 727 OD1 ASP A 107	40.647 0.738 -5.186 1.00 49.58 O
ANISOU 727 OD1 ASP A 107	6492 7299 5049 680 -90 325 O
ATOM 728 OD2 ASP A 107	39.803 -0.659 -3.734 1.00 57.05 O
ANISOU 728 OD2 ASP A 107	7368 8202 6108 595 -188 246 O
ATOM 729 N THR A 108	42.223 3.508 -1.208 1.00 37.74 N
ANISOU 729 N THR A 108	4893 5554 3894 638 111 479 N
ATOM 730 CA THR A 108	42.300 4.594 -0.220 1.00 39.10 C
ANISOU 730 CA THR A 108	5040 5672 4143 639 154 535 C
ATOM 731 C THR A 108	42.898 5.892 -0.800 1.00 40.49 C
ANISOU 731 C THR A 108	5254 5836 4293 680 231 610 C
ATOM 732 O THR A 108	42.332 6.997 -0.630 1.00 36.81 O
ANISOU 732 O THR A 108	4786 5365 3837 711 238 678 O
ATOM 733 CB THR A 108	43.079 4.134 1.056 1.00 34.93 C
ANISOU 733 CB THR A 108	4483 5070 3720 590 176 493 C
ATOM 734 OG1 THR A 108	42.357 3.058 1.676 1.00 31.89 O
ANISOU 734 OG1 THR A 108	4064 4693 3361 550 102 440 O
ATOM 735 CG2 THR A 108	43.210 5.248 2.081 1.00 33.69 C
ANISOU 735 CG2 THR A 108	4303 4857 3640 591 220 545 C
ATOM 736 N ALA A 109	44.027 5.750 -1.494 1.00 43.13 N
ANISOU 736 N ALA A 109	5626 6167 4595 681 291 597 N
ATOM 737 CA ALA A 109	44.610 6.863 -2.260 1.00 45.45 C
ANISOU 737 CA ALA A 109	5964 6460 4846 714 366 668 C
ATOM 738 C ALA A 109	43.629 7.552 -3.176 1.00 42.88 C
ANISOU 738 C ALA A 109	5669 6191 4431 766 335 732 C
ATOM 739 O ALA A 109	43.636 8.762 -3.177 1.00 44.81 O
ANISOU 739 O ALA A 109	5932 6409 4684 791 75 810 O
ATOM 740 CD ALA A 109	45.895 6.480 -3.037 1.00 43.35 C
ANISOU 740 CB ALA A 109	5730 6200 4540 708 432 639 C
ATOM 741 N LYS A 110	42.787 6.841 -3.936 1.00 45.79 N
ANISOU 741 N LYS A 110	6047 6635 4718 783 262 703 N
ATOM 742 CA LYS A 110	41.913 7.560 -4.846 1.00 49.53 C
ANISOU 742 CA LYS A 110	6562 7164 5102 837 232 769 C
ATOM 743 C LYS A 110	40.838 8.304 -4.113 1.00 54.16 C
ANISOU 743 C LYS A 110	7107 7738 5740 858 190 815 C
ATOM 744 O LYS A 110	40.475 9.390 -4.528 1.00 54.04 O
ISOU 744 O LYS A 110	7113 7728 5691 906 200 894 O
ATOM 745 CB LYS A 110	41.226 6.672 -5.882 1.00 56.99 C
ANISOU 745 CB LYS A 110	7514 8198 5941 853 158 727 C
ATOM 746 CG LYS A 110	42.094 5.687 -6.627 1.00 62.40 C

TABLE 3-continued

ANISOU	746	CG	LYS	A	110	8232	8909	6589	836	180	659	C
ATOM	747	CD	LYS	A	110	41.236	4.494	-7.045	1.00	65.75	C	
ANISOU	747	DD	LYS	A	110	8646	9397	6939	829	81	585	C
ATOM	748	CE	LYS	A	110	41.801	3.777	-8.272	1.00	71.49	C	
ANISOU	748	CE	LYS	A	110	9427	10177	7558	839	92	535	C
ATOM	749	NZ	LYS	A	110	40.734	2.917	-8.877	1.00	75.32	N	
ANISOU	749	NZ	LYS	A	110	9912	10736	7972	845	-15	483	N
ATOM	750	N	LEU	A	111	40.319	7.694	-3.038	1.00	51.15	N	
ANISOU	750	N	LEU	A	111	6658	7341	5436	822	142	765	N
ATOM	751	CA	LEU	A	111	39.269	8.241	-2.167	1.00	55.41	C	
ANISOU	751	CA	LEU	A	111	7147	7872	6033	835	101	792	C
ATOM	752	C	LEU	A	111	39.677	9.415	-1.294	1.00	51.59	C	
ANISOU	752	C	LEU	A	111	8659	7309	5632	843	164	846	C
ATOM	753	O	DEC	A	111	38.819	10.081	-0.794	1.00	44.12	O	
ANISOU	753	O	LEU	A	111	5684	6361	4717	871	138	879	O
ATOM	754	CB	LEU	A	111	38.835	7.174	-1.130	1.00	57.68	C	
ANISOU	754	CB	LEU	A	111	7371	8156	6387	780	49	718	C
ATOM	755	CG	LEU	A	111	38.105	5.876	-1.476	1.00	58.03	C	
ANISOU	755	CG	LEU	A	111	7394	8266	6390	754	-35	650	C
ATOM	756	CD1	LEU	A	111	37.794	5.159	-0.146	1.00	59.49	C	
ANISOU	756	CD1	LEU	A	111	7517	8421	6665	695	-64	600	C
ATOM	757	CD2	LEU	A	111	36.856	6.120	-2.321	1.00	51.51	C	
ANISOU	757	CD2	LEU	A	111	6562	7524	5484	802	-106	678	C
ATOM	758	N	LEU	A	112	40.966	9.624	-1.026	1.00	50.08	N	
ANISOU	758	N	LEU	A	112	6491	7053	5484	816	245	846	N
ATOM	759	CA	ILE	A	112	41.358	10.657	-0.047	1.00	49.44	C	
ANISOU	759	CA	ILE	A	112	6400	6891	5493	813	299	885	C
ATOM	760	C	ILE	A	112	40.824	12.030	-0.462	1.00	43.48	C	
ANISOU	760	C	ILE	A	112	5676	6130	4714	876	308	975	C
ATOM	761	O	ILE	A	112	40.923	12.417	-1.567	1.00	47.34	O	
ANISOU	761	O	ILE	A	112	6217	6646	5124	909	322	1022	O
ATOM	762	CB	ILE	A	112	42.889	10.668	0.277	1.00	50.17	C	
ANISOU	762	CB	ILE	A	112	6508	6917	5638	771	382	869	C
ATOM	763	CG1	ILE	A	112	43.150	9.751	1.475	1.00	58.24	C	
ANISOU	763	CG1	ILE	A	112	7478	7908	6741	716	365	794	C
ATOM	764	CG2	ILE	A	112	43.390	12.074	0.587	1.00	50.43	C	
ANISOU	764	CG2	ILE	A	112	6564	6878	5721	785	450	938	C
ATOM	765	CD1	ILE	A	112	44.498	9.062	1.403	1.00	61.79	C	
ANISOU	765	CD1	ILE	A	112	7937	8333	7206	675	413	746	C
ATOM	766	N	ASP	A	113	40.144	12.691	0.455	1.00	51.65	N	

TABLE 3-continued

ANISOU	766	N	ASP	A	113	6676	7134	5813	893	290	994	N
ATOM	767	CA	ASP	A	113	39.577	14.016	0.198	1.00	49.25	C	
ANISOU	767	CA	ASP	A	113	6399	6815	5500	958	293	1075	C
ATOM	768	C	ASP	A	113	40.673	15.004	0.560	1.00	44	50	C
ANISOU	768	C	ASP	A	113	5834	6117	4958	946	380	1116	C
ATOM	769	O	ASP	A	113	41.006	15.156	1.735	1.00	39.14	O	
ANISOU	769	O	ASP	A	113	5123	5377	4372	914	403	1089	O
ATOM	770	CB	ASP	A	113	38.355	14.176	1.111	1.00	49.83	C	
ANISOU	770	CB	ASP	A	113	6410	6902	5623	982	235	1063	C
ATOM	771	CG	ASP	A	113	37.907	15.621	1.249	1.00	54.99	C	
ANISOU	771	CG	ASP	A	113	7082	7510	6301	1047	246	1135	C
ATOM	772	OD1	ASP	A	113	38.359	16.496	0.517	1.00	52.05	O	
ANISOU	772	OD1	ASP	A	113	6775	7104	5897	1078	286	1204	O
ATOM	773	OD2	ASP	A	113	37.116	15.854	2.141	1.00	64.78	O	
ANISOU	773	OD2	ASP	A	113	8270	8748	7595	1064	217	1121	O
ATOM	774	N	ARG	A	114	41.257	15.629	-0.452	1.00	43.00	N	
ANISOU	774	N	ARG	A	114	5711	5915	4713	965	427	1178	N
ATOM	775	CA	ARG	A	114	42.451	16.424	-0.259	1.00	41.80	C	
ANISOU	775	CA	ARG	A	114	5595	5677	4611	939	516	1213	C
ATOM	776	C	ARG	A	114	42.135	17.756	0.441	1.00	40.22	C	
ANISOU	776	C	ARG	A	114	5402	5397	4481	972	528	1267	C
ATOM	777	O	ARG	A	114	43.035	18.309	1.113	1.00	39.62	O	
ANISOU	777	O	ARG	A	114	5332	5237	4485	936	590	1270	O
ATOM	778	CB	ARG	A	114	43.142	16.698	-1.601	1.00	45.18	C	
ANISOU	778	CB	ARG	A	114	6094	6121	4952	946	567	1269	C
ATOM	779	CG	ARG	A	114	44.149	15.608	-1.996	1.00	52.76	C	
ANISOU	779	CG	ARG	A	114	7049	7115	5881	893	602	1210	C
ATOM	780	CD	ARG	A	114	43.492	14.436	-2.723	1.00	57.95	C	
ANISOU	780	CD	ARG	A	114	7697	7874	6448	906	533	1162	C
ATOM	781	NE	ARG	A	114	44.417	13.401	-3.266	1.00	62.98	N	
ANISOU	781	NE	ARG	A	114	8339	8548	7043	866	565	1104	N
ATOM	782	CZ	ARG	A	114	45.624	13.695	-3.819	1.00	68.55	C	
ANISOU	782	CZ	ARG	A	114	9080	9236	7728	843	651	1123	C
ATOM	783	NH1	ARG	A	114	46.152	14.813	-3.925	1.00	61.99	N	
ANISOU	783	NH1	ARG	A	114	8288	8347	6917	846	720	1202	N
ATOM	784	NH2	ARG	A	114	46.340	12.577	-4.265	1.00	70.30	N	
ANISOU	784	NH2	ARG	A	114	9298	9499	7914	815	670	1059	N
ATOM	785	N	LYS	A	115	40.885	18.245	0.260	1.00	37.39	N	
ANISOU	785	N	LYS	A	115	5043	5067	4095	1040	468	1304	N
ATOM	786	CA	LYS	A	115	40.401	19.454	0.943	1.00	42.96	C	

TABLE 3-continued

ANISOU 786 CA LYS A 115	5752 5703 4867 1084 468 1346 C
ATOM 787 C LYS A 115	40.316 19.195 2.463 1.00 40.86 C
ANISOU 787 C LYS A 115	5418 5405 4703 1050 461 1275 C
ATOM 788 O LYS A 115	40.783 20.027 3.203 1.00 41.96 O
ANISOU 788 O LYS A 115	5567 5455 4919 1042 504 1288 O
ATOM 789 CB LYS A 115	39.016 19.973 0.427 1.00 44.68 C
ANISOU 789 CB LYS A 115	5976 5967 5034 1174 398 1395 C
ATOM 790 CG LYS A 115	38.962 20.370 -1.059 1.00 47.51 C
ANISOU 790 CG LYS A 115	6410 6357 5286 1220 395 1476 C
ATOM 791 CD LYS A 115	40.009 21.448 -1.375 1.00 51.11 C
ANISOU 791 CD LYS A 115	6945 6717 5756 1210 480 1551 C
ATOM 792 CE LYS A 115	39.534 22.485 -2.408 1.00 56.20 C
ANISOU 792 CE LYS A 115	7541 7227 6206 1287 466 1657 C
ATOM 793 NZ LYS A 115	40.532 23.619 -2.533 1.00 55.31 N
ANISOU 793 NZ LYS A 115	7632 7131 6254 1269 551 1733 N
ATOM 794 N SER A 116	39.751 18.058 2.923 1.00 41.32 N
ANISOU 794 N SER A 116	5409 5531 4760 1028 408 1203 N
ATOM 795 CA SER A 116	39.790 17.750 4.353 1.00 36.71 C
ANISOU 795 CA SER A 116	4767 4918 4265 988 408 1138 C
ATOM 796 C SER A 116	41.183 17.604 4.888 1.00 34.71 C
ANISOU 796 C SER A 116	4523 4597 4068 919 473 1109 C
ATOM 797 O SER A 116	41.445 18.019 5.993 1.00 35.48 O
ANISOU 797 O SER A 116	4603 4632 4247 900 495 1089 O
ATOM 798 CB SER A 116	39.047 16.438 4.787 1.00 39.22 C
ANISOU 798 CB SER A 116	5014 5316 4571 961 345 1065 C
ATOM 799 OG SER A 116	37.662 16.419 4.387 1.00 46.80 O
ANISOU 799 OG SER A 116	5946 6352 5483 1018 276 1079 O
ATOM 800 N LEU A 117	42.001 16.813 4.174 1.00 35.22 N
ANISOU 800 N LEU A 117	4605 4688 4088 878 495 1093 N
ATOM 801 CA LEU A 117	43.366 16.518 4.545 1.00 36.86 C
ANISOU 801 CA LEU A 117	4816 4847 4344 814 553 1060 C
ATOM 802 C LEU A 117	44.132 17.844 4.835 1.00 37.24 C
ANISOU 802 C LEU A 117	4900 4795 4453 812 070 1139 O
ATOM 803 O LEU A 117	44.809 17.938 5.903 1.00 33.93 O
ANISOU 803 O LEU A 117	4458 4316 4118 769 647 1072 O
ATOM 804 CB LEU A 117	44.052 15.738 3.404 1.00 34.97 C
ANISOU 804 CB LEU A 117	4602 4655 4030 792 572 1053 C
ATOM 805 CG LEU A 117	45.527 15.382 3.666 1.00 40.44 C
ANISOU 805 CG LEU A 117	5291 5306 4767 730 634 1017 C
ATOM 806 CD1 LEU A 117	45.736 14.542 4.939 1.00 35.91 C

TABLE 3-continued

ANISOU	806	CD1	LEU	A	117	4660	4717	4266	683	611	937	C
ATOM	807	CD2	LEU	A	117	46.166	14.709	2.447	1.00	40.38	C	
ANISOU	807	CD2	LEU	A	117	5312	5351	4681	719	659	1012	C
ATOM	808	N	ALA	A	118	44.054	18.821	3.906	1.00	36.70	N	
ANISOU	808	N	ALA	A	118	4892	4710	4343	854	644	1190	N
ATOM	809	CA	ALA	A	118	44.775	20.131	4.118	1.00	34.99	C	
ANISOU	809	CA	ALA	A	118	4717	4390	4186	848	708	1243	C
ATOM	810	C	ALA	A	118	44.305	20.827	5.396	1.00	35.57	C	
ANISOU	810	C	ALA	A	118	4766	4400	4348	864	691	1227	C
ATOM	811	O	ALA	A	118	45.126	21.302	6.198	1.00	32.87	O	
ANISOU	811	O	ALA	A	118	4423	3979	4087	824	734	1211	O
ATOM	812	CB	ALA	A	118	44.666	21.052	2.890	1.00	40.99	C	
ANISOU	812	CB	ALA	A	118	5552	5141	4862	894	781	1340	C
ATOM	813	N	ARG	A	119	42.983	20.823	5.624	1.00	33.64	N	
ANISOU	813	N	ARG	A	119	4497	4198	4086	922	627	1222	N
ATOM	814	CA	ARG	A	119	42.520	21.347	6.848	1.00	36.08	C	
ANISOU	814	CA	ARG	A	119	4776	4459	4472	937	612	1196	C
ATOM	815	C	ARG	A	119	43.021	20.605	8.089	1.00	33.17	C	
ANISOU	815	C	ARG	A	119	4353	4083	4168	873	615	1112	C
ATOM	816	O	ARG	A	119	43.318	21.240	9.096	1.00	31.95	O	
ANISOU	816	O	ARG	A	119	4194	3855	4089	861	636	1094	O
ATOM	817	CB	ARG	A	119	41.008	21.381	6.861	1.00	40.56	C	
ANISOU	817	CB	ARG	A	119	5315	5087	5008	1009	544	1201	C
ATOM	818	CG	ARG	A	119	40.519	22.605	6.155	1.00	39.12	C	
ANISOU	818	CG	ARG	A	119	5189	4868	4805	1085	543	1284	C
ATOM	819	CD	ARG	A	119	39.086	22.837	6.590	1.00	43.00	C	
ANISOU	819	CD	ARG	A	119	5639	5398	5301	1158	481	1274	C
ATOM	820	NE	ARG	A	119	38.262	21.597	6.498	1.00	39.39	N	
ANISOU	820	NE	ARG	A	119	5116	5059	4792	1153	421	1227	N
ATOM	821	CZ	ARG	A	119	37.707	21.134	5.399	1.00	36.71	C	
ANISOU	821	CZ	ARG	A	119	4781	4801	4366	1182	380	1252	C
ATOM	822	NH1	ARG	A	119	37.885	21.765	4.210	1.00	38.98	N	
ANISOU	822	NH1	ARG	A	119	5140	5074	4597	1220	391	1330	N
ATOM	823	NH2	ARG	A	119	36.974	20.036	5.473	1.00	36.03	N	
ANISOU	823	NH2	ARG	A	119	4630	4811	4247	1169	325	1201	N
ATOM	824	N	ILE	A	120	43.058	19.268	8.021	1.00	32.10	N	
ANISOU	824	N	ILE	A	120	4178	4021	3997	836	589	1061	N
ATOM	825	CA	ILE	A	120	43.491	18.464	9.175	1.00	32.96	C	
ANISOU	825	CA	ILE	A	120	4238	4125	4159	777	585	985	C
ATOM	826	C	ILE	A	120	44.995	18.764	9.502	1.00	33.48	C	

TABLE 3-continued

ANISOU 826 C ILE A 120	4323 4112 4285 721 649 975 C
ATOM 827 O ILE A 120	45.400 18.895 10.657 1.00 31.27 O
ANISOU 827 O 1E A 120	4022 3784 4077 690 658 934 O
ATOM 828 CB ILE A 120	43.242 16.950 8.905 1.00 32.22 C
ANISOU 828 CB ILE A 120	4108 4120 4013 750 542 937 C
ATOM 829 CG1 ILE A 120	41.711 16.714 8.754 1.00 85.60 C
ANISOU 829 CG1 ILE A 120	4507 4625 4395 799 476 941 C
ATOM 830 CG2 ILE A 120	43.872 16.110 10.017 1.00 34.50 C
ANISOU 830 CG2 ILE A 120	4358 4394 4355 687 541 866 C
ATOM 831 CD1 ILE A 120	41.307 15.314 8.268 1.00 29.54 C
ANISOU 831 CD1 ILE A 120	3712 3946 3567 778 427 904 C
ATOM 832 N VAL A 121	45.810 18.866 8.472 1.00 32.18 N
ANISOU 832 N VAL A 121	4197 3940 4090 707 692 1012 N
ATOM 833 CA VAL A 121	47.216 19.275 8.636 1.00 31.34 C
ANISOU 833 CA VAL A 121	4106 3763 4040 655 757 1011 C
ATOM 834 C VAL A 121	47.378 20.538 9.410 1.00 30.32 C
ANISOU 834 C VAL A 121	3994 3538 3987 659 783 1030 C
ATOM 835 O VAL A 121	43.197 20.605 10.356 1.00 31.39 O
ANISOU 835 O VAL A 121	4109 3621 4197 611 803 987 O
ATOM 836 CB VAL A 121	47.855 19.555 7.290 1.00 33.75 C
ANISOU 836 CB VAL A 121	4459 4071 4293 653 807 1070 C
ATOM 837 CG1 VAL A 121	49.246 20.239 7.517 1.00 30.53 C
ANISOU 837 CG1 VAL A 121	4065 3581 3955 600 881 1079 C
ATOM 838 CG2 VAL A 121	47.976 18.227 6.482 1.00 36.71 C
ANISOU 838 CG2 VAL A 121	4818 4537 4593 641 792 1040 C
ATOM 839 N GLU A 122	46.557 21.550 9.081 1.00 32.23 N
ANISOU 839 N GLU A 122	4274 3756 4215 719 775 1090 N
ATOM 840 CA GLU A 122	46.653 22.817 9.771 1.00 34.78 C
ANISOU 840 CA GLU A 122	4622 3981 4612 730 797 1108 C
ATOM 841 C GLU A 122	46.246 22.699 11.230 1.00 33.87 C
ANISOU 841 C GLU A 122	4460 3853 4555 728 763 1039 C
ATOM 842 O GLU A 122	46.911 23.237 12.098 1.00 31.48 O
ANISOU 842 O GLU A 122	4159 3475 4328 695 788 1014 O
ATOM 843 CB GLU A 122	45.742 23.831 9.037 1.00 44.09 C
ANISOU 843 CB GLU A 122	5853 5143 5756 806 786 1186 C
ATOM 844 CG GLU A 122	46.165 25.275 9.084 1.00 49.83 C
ANISOU 844 CG GLU A 122	6638 5755 6540 813 830 1239 C
ATOM 845 CD GLU A 122	47.662 25.462 9.009 1.00 58.44 C
ANISOU 845 CD GLU A 122	7744 6785 7674 733 897 1242 C
ATOM 846 OE1 GLU A 122	48.343 24.809 8.145 1.00 47.68 O

TABLE 3-continued

ANISOU	846	OE1	GLU	A	122	6383	5471	6263	696	927	1255	O
ATOM	847	OE2	GLU	A	122	48.150	26.282	9.859	1.00	67.06	0	
ANISOU	847	OE2	GLU	A	122	8846	7781	8853	709	920	1227	O
ATOM	848	N	ARG	A	123	45.167	21.971	11.503	1.00	35.91	N	
ANISOU	848	N	ARG	A	123	4678	4189	4776	759	707	1008	N
ATOM	849	CA	ARG	A	123	44.795	21.667	12.922	1.00	35.90	C	
ANISOU	849	CA	ARG	A	123	4628	4191	4820	749	677	939	C
ATOM	850	C	ARG	A	123	45.938	21.018	13.729	1.00	34.25	C	
ANISOU	850	C	ARG	A	123	4393	3962	4659	672	694	878	C
ATOM	851	O	ARG	A	123	46.230	21.434	14.843	1.00	37.66	O	
ANISOU	851	O	ARG	A	123	4816	4339	5153	654	701	841	O
ATOM	852	CB	ARG	A	123	43.550	20.813	12.917	1.00	34.52	C	
ANISOU	852	CB	ARG	A	123	4410	4115	4590	781	620	920	C
ATOM	853	CG	ARG	A	123	42.365	21.652	12.351	1.00	37.48	C	
ANISOU	853	CG	ARG	A	123	4804	4503	4935	867	598	974	C
ATOM	854	CD	ARG	A	123	41.006	21.129	12.762	1.00	45.89	C	
ANISOU	854	CD	ARG	A	123	5814	5652	5972	904	542	945	C
ATOM	855	NE	ARG	A	123	39.856	21.799	12.105	1.00	46.24	N	
ANISOU	855	NE	ARG	A	123	5867	5722	5981	990	514	995	N
ATOM	856	CZ	ARG	A	123	39.122	22.764	12.660	1.00	53.34	C	
ANISOU	856	CZ	ARG	A	123	6763	6589	6914	1053	507	1000	C
ATOM	857	NH1	ARG	A	123	39.437	23.260	13.868	1.00	41.17	N	
ANISOU	857	NH1	ARG	A	123	5217	4984	5443	1038	530	959	N
ATOM	858	NH2	ARG	A	123	38.047	23.228	12.002	1.00	53.51	N	
ANISOU	858	NH2	ARG	A	123	6786	6647	6900	1134	473	1043	N
ATOM	859	N	VAL	A	124	46.565	19.998	13.166	1.00	32.56	N	
ANISOU	859	N	VAL	A	124	4167	3793	4412	631	698	866	N
ATOM	860	CA	VAL	A	124	47.708	19.369	13.800	1.00	32.16	C	
ANISOU	860	CA	VAL	A	124	4092	3724	4405	564	712	812	C
ATOM	861	O	VAL	A	124	46.859	20.353	14.048	1.00	33.36	C	
ANISOU	861	O	VAL	A	124	4268	3781	4627	531	766	821	C
ATOM	862	O	VAL	A	124	49.429	20.433	15.159	1.00	37.71	O	
ANISOU	862	O	VAL	A	124	4799	4288	5241	495	766	772	O
ATOM	863	CB	VAL	A	124	48.180	18.162	13.011	1.00	32.66	C	
ANISOU	863	CB	VAL	A	124	4142	3849	4420	536	709	799	C
ATOM	864	CG1	VAL	A	124	49.434	17.545	13.702	1.00	32.71	C	
ANISOU	864	CG1	VAL	A	124	4120	3830	4477	472	722	742	C
ATOM	865	CG2	VAL	A	124	47.027	17	127	12.969	1.00	29.93	C
ANISOU	865	CG2	VAL	A	124	3768	3590	4015	558	649	779	C
ATOM	866	N	HIS	A	125	49.110	21.163	13.064	1.00	34.73	N	

TABLE 3-continued

ANISOU 866 N HIS A 125	4485 3921 4789 545 807 885 N
ATOM 867 CA HIS A 125	50.181 22.136 13.169 1.00 36.69 C
ANISOU 867 CA HIS A 125	4758 4078 5103 508 661 901 C
ATOM 868 C HIS A 125	49.905 23.062 14.321 1.00 35.30 C
ANISOU 868 C HIS A 125	4589 3829 4994 520 851 881 C
ATOM 869 O HIS A 125	50.807 23.304 15.133 1.00 36.67 O
ANISOU 869 O HIS A 125	4749 3946 5237 471 867 841 O
ATOM 870 CB HIS A 125	50.321 22.923 11.881 1.00 33.31 C
ANISOU 870 CB HIS A 125	4385 3627 4644 526 906 985 C
ATOM 871 CG HIS A 125	51.330 24.030 12.001 1.00 38.67 C
ANISOU 871 CG HIS A 125	5093 4204 5395 485 933 1009 C
ATOM 872 ND1 HIS A 125	52.611 23.794 12.261 1.00 39.87 N
ANISOU 872 ND1 HIS A 125	5218 4334 5565 415 998 975 N
ATOM 873 CD2 HIS A 125	51.179 25.429 12.004 1.00 45.43 C
ANISOU 873 CD2 HIS A 125	6003 4968 6290 505 985 1060 C
ATOM 874 CE1 HIS A 125	53.281 24.971 12.370 1.00 40.49 C
ANISOU 874 CE1 HIS A 125	5328 4315 5740 385 1043 1004 C
ATOM 875 NE2 HIS A 125	52.418 25.961 12.180 1.00 41.10 N
ANISOU 875 NE2 HIS A 125	5459 4346 5810 439 1036 1058 N
ATOM 876 N GLN A 126	48.656 23.527 14.438 1.00 32.05 N
ANISOU 876 N GLN A 126	4191 3424 4561 587 819 900 N
ATOM 877 CA GLN A 126	48.274 24.457 15.521 1.00 38.06 C
ANISOU 877 CA GLN A 126	4961 4118 5381 610 809 877 C
ATOM 878 C GLN A 126	48.218 23.779 16.908 1.00 39.80 C
ANISOU 878 C GLN A 126	5131 4363 5627 586 775 792 C
ATOM 879 O GLN A 126	48.347 24.450 17.949 1.00 37.09 O
ANISOU 879 O GLN A 126	4792 3957 5342 581 775 756 O
ATOM 880 CB GLN A 126	46.901 25.130 15.264 1.00 35.18 C
ANISOU 880 CB GLN A 126	4621 3760 4986 697 784 917 C
ATOM 881 CG GLN A 126	46.800 25.972 13.963 1.00 49.72 C
ANISOU 881 CG GLN A 126	6523 5567 6601 734 811 1008 C
ATOM 882 CD GLN A 126	47.814 27.137 13.894 1.00 54.30 C
ANISOU 882 CD GLN A 126	7156 6027 7447 700 864 1041 C
ATOM 883 OE1 GLN A 126	48.211 27.690 14.920 1.00 58.09 O
ANISOU 883 OE1 GLN A 126	7635 6434 8001 677 870 997 O
ATOM 884 NE2 GLN A 126	48.238 27.501 12.674 1.00 60.30 N
ANISOU 884 NE2 GLN A 126	7964 6769 8180 694 902 1117 N
ATOM 885 N ALA A 127	48.030 22.456 16.930 1.00 37.37 N
ANISOU 885 N ALA A 127	4781 4144 5274 569 745 761 N
ATOM 886 CA ALA A 127	47.625 21.836 18.169 1.00 35.09 C

TABLE 3-continued

ANISOU 886 CA ALA A 127	4451 3890 4993 561 706 696 C
ATOM 887 C ALA A 127	48.787 21.844 19.165 1.00 37.25 C
ANISOU 887 C ALA A 127	4700 4096 5318 500 717 642 C
ATOM 888 O ALA A 127	49.941 21.603 18.802 1.00 40.24 O
ANISOU 888 O ALA A 127	5091 4468 5731 450 742 641 O
ATOM 889 CB ALA A 127	47.113 20.441 17.905 1.00 37.55 C
ANISOU 889 CB ALA A 127	4725 4301 5241 556 669 682 C
ATOM 890 N GLU A 128	48.506 22.146 20.420 1.00 33.40 N
ANISOU 890 N GLU A 128	4217 3600 4875 504 698 594 N
ATOM 891 CA GLU A 128	49.531 22.004 21.444 1.00 36.86 C
ANISOU 891 CA GLU A 128	4640 4000 5365 448 697 536 C
ATOM 892 C GLU A 128	49.523 20.579 22.041 1.00 35.36 C
ANISOU 892 C GLU A 128	4408 3684 5144 417 657 491 C
ATOM 893 O GLU A 128	50.546 20.049 22.531 1.00 32.86 O
ANISOU 893 O GLU A 128	4074 3556 4856 365 650 451 O
ATOM 894 CB GLU A 128	49.284 23.046 22.531 1.00 43.74 C
ANISOU 894 CB GLU A 128	5529 4809 6282 467 695 504 C
ATOM 895 CG GLU A 128	49.786 24.436 22.054 1.00 53.11 C
ANISOU 895 CG GLU A 128	6763 5895 7523 472 737 540 C
ATOM 896 CD GLU A 128	49.519 25.539 23.078 1.00 61.91 C
ANISOU 896 CD GLU A 128	7901 6938 8685 496 734 505 C
ATOM 897 OE1 GLU A 128	48.515 25.448 23.804 1.00 62.23 O
ANISOU 897 OE1 GLU A 128	7928 7018 8699 539 707 475 O
ATOM 898 OE2 GLU A 128	50.299 26.518 23.162 1.00 73.05 O
ANISOU 898 OE2 GLU A 128	9343 8252 10160 472 760 505 O
ATOM 899 N PHE A 129	48.361 19.919 21.952 1.00 31.26 N
ANISOU 899 N PHE A 129	3872 3441 4566 449 628 497 N
ATOM 900 CA PHE A 129	48.305 18.561 22.588 1.00 35.38 C
ANISOU 900 CA PHE A 129	4359 4027 5058 416 588 457 C
ATOM 901 C PHE A 129	47.368 17.746 21.742 1.00 37.44 C
ANISOU 901 C PHE A 129	4605 4367 5252 439 569 487 C
ATOM 902 O PHE A 129	46.319 18.272 21.274 1.00 35.24 O
ANISOU 902 O PHE A 129	4333 4111 4947 492 571 521 O
ATOM 903 CB PHE A 129	47.783 18.636 24.046 1.00 33.63 C
ANISOU 903 CB PHE A 129	4125 3812 4842 418 564 409 C
ATOM 904 CG PHE A 129	47.468 17.243 24.657 1.00 34.07 C
ANISOU 904 CG PHE A 129	4150 3938 4856 388 523 380 C
ATOM 905 CD1 PHE A 129	48.457 16.521 25.281 1.00 36.32 C
ANISOU 905 CD1 PHE A 129	4428 4211 5161 337 504 342 C
ATOM 906 CD2 PHE A 129	46.178 16.677 24.528 1.00 41.60 C

TABLE 3-continued

ANISOU	906	CD2	PHE	A	129	5083	4969	5754	412	501	394	C
ATOM	907	CE1	PHE	A	129	48.153	15.263	25.870	1.00	39.68	C	
ANISOU	907	CE1	PHE	A	129	4834	4694	5550	309	463	320	C
ATOM	908	CE2	PHE	A	129	45.902	15.433	25.073	1.00	37.79	C	
ANISOU	908	CE2	PHE	A	129	4578	4544	5238	378	465	371	C
ATOM	909	CZ	PHE	A	129	46.887	14.744	25.746	1.00	32.27	C	
ANISOU	909	CZ	PHE	A	129	3879	3825	4557	328	447	337	C
ATOM	910	N	ILE	A	130	47.762	16.501	21.479	1.00	31.98	N	
ANISOU	910	N	ILE	A	130	3897	3717	4538	403	548	474	N
ATOM	911	CA	ILE	A	130	46.958	15.667	20.616	1.00	31.17	C	
ANISOU	911	CA	IEL	A	130	3783	3686	4374	418	526	498	C
ATOM	912	C	ILE	A	130	46.658	14.374	21.358	1.00	30.70	C	
ANISOU	912	C	ILE	A	130	3695	3677	4290	384	481	461	C
ATOM	913	O	ILE	A	130	47.603	13.585	21.758	1.00	28.44	O	
ANISOU	913	O	ILE	A	130	3405	3377	4023	339	468	428	O
ATOM	914	CB	ILE	A	130	47.696	15.372	19.271	1.00	33.45	C	
ANISOU	914	C8	ILE	A	130	4086	3976	4649	410	546	526	C
ATOM	915	CG1	ILE	A	130	47.938	16.704	18.504	1.00	30.29	C	
ANISOU	915	CG1	ILE	A	130	3719	3524	4267	441	594	573	C
ATOM	916	CG2	ILE	A	130	46.860	14.412	18.365	1.00	31.42	C	
ANISOU	916	CG2	ILE	A	130	3819	3797	4323	424	517	543	C
ATOM	917	CD1	ILL	A	130	48.794	16.547	17.255	1.00	32.50	C	
ANISOU	917	CD1	ILE	A	130	4015	3800	4534	429	625	601	C
ATOM	918	N	GLY	A	131	45.373	14.125	21.519	1.00	27.60	N	
ANISOU	918	N	GLY	A	131	3284	3344	3857	405	457	467	N
ATOM	919	CA	GLY	A	131	44.900	12.910	22.209	1.00	27.55	C	
ANISOU	919	CA	GLY	A	131	3253	3390	3824	370	416	439	C
ATOM	920	C	GLY	A	131	44.410	11.939	21.146	1.00	29.39	C	
ANISOU	920	C	GLY	A	131	3477	3681	4008	367	390	457	C
ATOM	921	O	GLY	A	131	43.668	12.341	20.210	1.00	28.36	O	
ANISOU	921	O	GLY	A	131	3346	3582	3848	409	396	492	O
ATOM	922	N	CYS	A	132	44.734	10.645	21.272	1.00	26.18	N	
ANISOU	922	N	CYS	A	132	3066	3291	3589	322	358	434	N
ATOM	923	CA	CYS	A	132	44.253	9.689	20.251	1.00	29.13	C	
ANISOU	923	CA	CYS	A	132	3434	3717	3916	318	330	445	C
ATOM	924	C	CYS	A	132	43.496	8.547	20.939	1.00	28.79	C	
ANISOU	924	C	CYS	A	132	3369	3721	3848	278	285	426	C
ATOM	925	O	CYS	A	132	43.882	8.187	22.045	1.00	26.72	O	
ANISOU	925	O	CYS	A	132	3108	3436	3607	243	274	400	O
ATOM	926	CB	CYS	A	132	45.420	9.062	19.488	1.00	29.60	C	

TABLE 3-continued

ANISOU 926	CB	CYS	A	132	3514	3750	3982	300	332	434	C	
ATOM	927	SG	CYS	A	132	46.470	10.328	18.719	1.00	30.93	S	
ANISOU 927	SG	CYS	A	132	3707	3862	4182	332	392	458	S	
ATOM	928	N	ILE	A	133	42.435	8.042	25.306	1.00	24.86	N	
ANISOU 928	N	ILE	A	133	2852	3286	3306	282	258	440	N	
ATOM	929	CA	ILE	A	133	41.550	7.081	20.979	1.00	30.37	C	
ANISOU 929	CA	ILE	A	133	3525	4033	3982	241	220	427	C	
ATOM	930	C	ILE	A	133	41.246	5.958	20.000	1.00	29.47	C	
ANISOU 930	C	ILE	A	133	3411	3955	3832	220	180	426	C	
ATOM	931	O	ILE	A	133	40	842	6.213	18.866	1.00	30.82	O
ANISOU 931	O	ILE	A	133	3579	4156	3974	255	180	445	O	
ATOM	932	CB	ILE	A	133	40.203	7.712	21.503	1.00	33.09	C	
ANISOU 932	CB	ILE	A	133	3829	4431	4311	263	226	441	C	
ATOM	933	CG1	ILE	A	133	40.376	8.957	22.358	1.00	32.51	C	
ANISOU 933	CG1	ILE	A	133	3758	4324	4270	295	266	439	C	
ATOM	934	CG2	ILE	A	133	39.403	6.680	22.288	1.00	32.09	C	
ANISOU 934	CG2	ILE	A	133	3675	4354	4164	208	192	428	C	
ATOM	935	CD1	ILE	A	133	39.039	9.717	22.376	1.00	42.57	C	
ANISOU 935	CD1	ILE	A	133	4992	5655	5526	341	275	456	C	
ATOM	936	N	GLY	A	134	41.410	4.689	20.459	1.00	27.55	N	
ANISOU 936	N	GLY	A	134	3174	3707	3587	163	143	403	N	
ATOM	937	CA	GLY	A	134	41.019	3.566	19.654	1.00	29.95	C	
ANISOU 937	CA	GLY	A	134	3479	4043	3859	138	100	396	C	
ATOM	938	C	GLY	A	134	41.244	2.272	20.403	1.00	31.31	C	
ANISOU 938	C	GLY	A	134	3664	4193	4038	74	61	372	C	
ATOM	939	O	GLY	A	134	42.119	2.213	21.312	1.00	30.51	O	
ANISOU 939	O	GLY	A	134	3582	4040	3970	57	67	359	O	
ATOM	940	N	VAL	A	135	40.405	1.276	20.085	1.00	29.72	N	
ANISOU 940	N	VAL	A	135	3452	4033	3809	36	17	369	N	
ATOM	941	CA	VAL	A	135	40.516	-0.046	20.722	1.00	26.74	C	
ANISOU 941	CA	VAL	A	135	3091	3632	3436	-29	-20	351	C	
ATOM	942	C	VAL	A	135	40.655	-1.205	19.812	1.00	26.12	C	
ANISOU 942	C	VAL	A	135	3037	3544	3343	-51	-71	330	C	
ATOM	943	O	VAL	A	135	40.426	-1.110	18.573	1.00	26.71	O	
ANISOU 943	O	VAL	A	135	3108	3648	3391	-20	-75	328	O	
ATOM	944	CB	VAL	A	135	39.316	-0.324	21.695	1.00	31.15	C	
ANISOU 944	CB	VAL	A	135	3615	4239	3981	-79	-40	366	C	
ATOM	945	CG1	VAL	A	135	39.198	0.773	22.808	1.00	30.31	C	
ANISOU 945	CG1	VAL	A	135	3488	4140	3887	-59	5	380	C	
ATOM	946	CG2	VAL	A	135	38.021	-0.445	20.904	1.00	31.98	C	

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TABLE 3-continued

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ANISOU 946 CG2 VAL A 135	3679 4420 4051 -82 -59 376 C
ATOM 947 N GLY A 136	41.086 -2.357 20.349 1.00 26.13 N
ANISOU 947 N GLY A 136	3069 3500 3359 -101 -109 312 N
ATOM 948 CA GLY A 136	41.239 -3.561 19.501 1.00 28.01 C
ANISOU 948 CA GLY A 136	3336 3719 3587 -122 -157 284 C
ATOM 949 C GLY A 136	42.232 -3.238 18.389 1.00 28.42 C
ANISOU 949 C GLY A 136	3409 3749 3540 -64 -136 264 C
ATOM 950 O GLY A 136	43.312 -2.695 18.661 1.00 30.81 O
ANISOU 950 O GLY A 136	3724 4009 3972 -32 -102 260 O
ATOM 951 N ALA A 137	41.917 -3.588 17.143 1.00 27.12 N
ANISOU 951 N ALA A 137	3249 3614 3442 -50 -156 250 N
ATOM 952 CA ALA A 137	42.858 -3.364 16.052 1.00 28.11 C
ANISOU 952 CA ALA A 137	3397 3724 3561 2 -133 230 C
ATOM 953 C ALA A 137	43.092 -1.885 15.789 1.00 27.59 C
ANISOU 953 C ALA A 137	3313 3678 3493 59 -70 259 C
ATOM 954 O ALA A 137	44.141 -1.505 15.229 1.00 28.79 O
ANISOU 954 O ALA A 137	3482 3803 3653 98 -35 249 O
ATOM 955 CB ALA A 137	42.398 -3.994 14.764 1.00 24.83 C
ANISOU 955 CB ALA A 137	2892 3343 3100 6 -157 208 C
ATOM 956 N SER A 138	42.115 -1.060 16.126 1.00 25.42 N
ANISOU 956 N SER A 138	3003 3449 3207 64 -55 294 N
ATOM 957 CA SER A 138	42.319 0.416 15.951 1.00 25.59 C
ANISOU 957 CA SER A 138	3025 3491 3245 119 4 325 C
ATOM 958 C SER A 138	43.365 0 954 16.905 1 00 27.30 C
ANISOU 958 C SER A 138	3237 3636 3500 123 41 324 C
ATOM 959 O SER A 138	43.903 2.033 16.653 1.00 28.54 O
ANISOU 959 O SER A 138	3395 3779 3668 164 90 341 O
ATOM 960 CB SER A 138	40.979 1.196 16.114 1.00 23.36 C
ANISOU 960 CB SER A 138	2691 3261 2932 131 9 360 C
ATOM 961 OG SER A 138	40.287 1.095 14.815 1.00 27.41 O
ANISOU 961 OG SER A 138	3196 3827 3391 154 -13 365 O
ATOM 962 N SER A 139	43.594 0.265 18.039 1.00 28.06 N
ANISOU 962 N SER A 139	3338 3697 3625 78 16 308 N
ATOM 963 CA SER A 139	44.801 0.623 18.900 1.00 27.74 C
ANISOU 963 CA SER A 139	3310 3595 3634 81 42 298 C
ATOM 964 C SER A 139	46.105 0.724 18.043 1.00 30.32 C
ANISOU 964 C SER A 139	2657 3887 3976 115 68 278 C
ATOM 965 O SER A 139	46.939 1.606 18.260 1.00 26.07 O
ANISOU 965 O SER A 139	3116 3319 3470 138 111 282 O
ATOM 966 CB SER A 139	45.052 -0.470 19.955 1.00 31.68 C

TABLE 3-continued

ANISOU 966 CB SER A 139	3825 4058 4154 31 -2 277 C
ATOM 967 OG SER A 139	46.224 -0.214 20.710 1.00 35.53 O
ANISOU 967 OG SER A 139	4323 4490 4685 36 13 264 O
ATOM 968 N ILE A 140	46.267 -0.189 17.103 1.00 28.39 N
ANISOU 968 N ILE A 140	3430 3648 3710 115 43 254 N
ATOM 969 CA ILE A 140	47.532 -0.290 16.273 1.00 32.68 C
ANISOU 969 CA ILE A 140	3990 4163 4264 145 67 227 C
ATOM 970 C ILE A 140	47.662 0.974 15.448 1.00 30.47 C
ANISOU 970 C ILE A 140	3703 3907 3969 189 127 257 C
ATOM 971 O ILE A 140	48.760 1.564 15.357 1.00 26.83 O
ANISOU 971 O ILE A 140	3242 3415 3539 208 173 253 O
ATOM 972 CB ILE A 140	47.468 -1.551 15.329 1.00 30.71 C
ANISOU 972 CB ILE A 140	3764 3924 3982 140 24 192 C
ATOM 973 CG1 ILE A 140	47.781 -2.853 16.111 1.00 30.06 C
ANISOU 973 CG1 ILE A 140	3699 3793 3928 102 -32 157 C
ATOM 974 CG2 ILE A 140	48.426 -1.466 14.130 1.00 36.56 C
ANISOU 974 CG2 ILE A 140	4517 4665 4711 181 60 170 C
ATOM 975 CD1 ILE A 140	46.904 -3.159 17.312 1.00 31.87 C
ANISOU 975 CD1 ILE A 140	3923 4021 4165 52 -70 177 C
ATOM 976 N VAL A 141	46.531 1.414 14.876 1.00 28.15 N
ANISOU 976 N VAL A 141	3401 3665 3629 202 127 288 N
ATOM 977 CA VAL A 141	46.531 2.644 14.054 1.00 27.28 C
ANISOU 977 CA VAL A 141	3291 3577 3499 247 180 324 C
ATOM 978 C VAL A 141	46.780 3.927 14.864 1.00 26.91 C
ANISOU 978 C VAL A 141	3231 3499 3494 258 225 353 C
ATOM 979 O VAL A 141	47.613 4.759 14.418 1.00 26.46 O
ANISOU 979 O VAL A 141	3182 3420 3452 282 278 367 O
ATOM 980 CB VAL A 141	45.226 2.772 13.169 1.00 28.79 C
ANISOU 980 CB VAL A 141	3477 3834 3626 266 160 351 C
ATOM 981 CG1 VAL A 141	45.233 4.083 12.344 1.00 30.12 C
ANISOU 981 CG1 VAL A 141	3653 4020 3773 316 213 396 C
ATOM 982 CG2 VAL A 141	45.152 1.518 12.292 1.00 27.71 C
ANISOU 982 CG2 VAL A 141	3359 3722 3446 255 117 314 C
ATOM 983 N GLY A 142	46.088 4.053 16.010 1.00 27.16 N
ANISOU 983 N GLY A 142	3244 3531 3545 237 206 360 N
ATOM 984 CA GLY A 142	46.241 5.176 16.920 1.00 27.43 C
ANISOU 984 CA GLY A 142	3268 3534 3619 245 240 378 C
ATOM 985 C GLY A 142	47.685 5.234 17.497 1.00 31.29 C
ANISOU 985 C GLY A 142	3763 3961 4163 232 262 352 C
ATOM 986 O GLY A 142	48.285 6.285 17.578 1.00 27.91 O

TABLE 3-continued

ANISOU 986 O GLY A 142	3336 3502 3767 248 307 365 O
ATOM 987 N ARG A 143	43.279 4.079 17.832 1.00 26.29 N
ANISOU 987 N ARG A 143	3136 3308 3545 202 226 314 N
ATOM 988 CA ARG A 143	49.683 4.041 18.237 1.00 27.35 C
ANISOU 988 CA ARG A 143	3271 3369 3730 195 241 286 C
ATOM 989 C ARG A 143	50.594 4.496 17.127 1.00 25.76 C
ANISOU 989 C ARG A 143	3073 3182 3533 222 290 288 C
ATOM 990 O ARG A 143	51.586 5.148 17.393 1.00 30.35 O
ANISOU 990 O ARG A 143	3646 3726 4160 223 326 283 O
ATOM 991 CB ARG A 143	50.086 2.594 18.629 1.00 29.70 C
ANISOU 991 CB ARG A 143	3577 3669 4037 167 187 245 C
ATOM 992 CG ARG A 143	49.574 2.098 20.004 1.00 33.21 C
ANISOU 992 CG ARG A 143	4023 4105 4492 130 141 241 C
ATOM 993 CD ARG A 143	49.904 3.023 21.222 1.00 43.54 C
ANISOU 993 CD ARG A 143	5321 5385 5839 125 160 246 C
ATOM 994 NE ARG A 143	49.194 2.663 22.511 1.00 46.52 N
ANISOU 994 NE ARG A 143	5700 5766 6208 91 123 249 N
ATOM 995 CZ ARG A 143	49.402 3.206 23.728 1.00 42.15 C
ANISOU 995 CZ ARG A 143	5144 5192 5680 81 126 246 C
ATOM 996 NH1 ARG A 143	50.266 4.220 23.885 1.00 39.65 N
ANISOU 996 NH1 ARG A 143	4820 4843 5404 100 164 239 N
ATOM 997 NH2 ARG A 143	48.744 2.716 24.815 1.00 52.62 N
ANISOU 997 NH2 ARG A 143	6476 6528 6988 48 92 249 N
ATOM 998 N TYR A 144	50.335 4.084 15.907 1.00 25.18 N
ANISOU 998 N TYR A 144	3012 3145 3412 239 291 291 N
ATOM 999 CA TYR A 144	51.226 4.573 14.831 1.00 26.37 C
ANISOU 999 CA TYR A 144	3166 3294 3559 264 346 296 C
ATOM 1000 C TYR A 144	51.129 6.082 14.688 1.00 28.45 C
ANISOU 1000 C TYR A 144	3430 3551 3830 282 402 344 C
ATOM 1001 O TYR A 144	52.116 6.752 14.455 1.00 26.48 O
ANISOU 1001 O TYR A 144	3177 3274 3612 286 453 349 O
ATOM 1002 CB TYR A 144	50.929 3.882 13.489 1.00 24.48 C
ANISOU 1002 CB TYR A 144	2944 3101 3256 282 338 290 C
ATOM 1003 CG TYR A 144	51.985 4.190 12.423 1.00 26.69 C
ANISOU 1003 CG TYR A 144	3229 3384 3529 304 396 287 C
ATOM 1004 CD1 TYR A 144	53.398 3.902 12.643 1.00 27.05 C
ANISOU 1004 CD1 TYR A 144	3257 3393 3626 296 418 248 C
ATOM 1005 CD2 TYR A 144	51.568 4.676 11.162 1.00 30.26 C
ANISOU 1005 CD2 TYR A 144	3700 3879 3919 332 428 322 C
ATOM 1006 CE1 TYR A 144	54.339 4.159 11.633 1.00 31.03 C

TABLE 3-continued

ANISOU	1006	CE1	TYR	A	144	3760	3909	4122	313	478	244	C
ATOM	1007	CE2	TYR	A	144	52.531	4.908	10.176	1.00	31.11	C	
ANISOU	1007	CE2	TYR	A	144	3813	3994	4012	348	486	322	C
ATOM	1008	CZ	TYR	A	144	53.890	4.646	10.432	1.00	29.62	C	
ANISOU	1008	CZ	TYR	A	144	3604	3773	3877	337	512	282	C
ATOM	1009	OH	TYR	A	144	54.769	4.941	9.426	1.00	30.64	O	
ANISOU	1009	OH	TYR	A	144	3734	3919	3989	352	577	285	O
ATOM	1010	N	LEU	A	145	49.918	6.614	14.796	1.00	28.11	N	
ANISOU	1010	N	LEU	A	145	3390	3531	3759	294	391	379	N
ATOM	1011	CA	LEU	A	145	49.761	8.086	14.758	1.00	24.42	C	
ANISOU	1011	CA	LEU	A	145	2927	3047	3304	316	438	425	C
ATOM	1012	C	LEU	A	145	50.525	8.820	15.892	1.00	25.57	C	
ANISOU	1012	C	LEU	A	145	3061	3132	3522	298	461	416	C
ATOM	1013	O	LEU	A	145	51.256	9.816	15.606	1.00	27.08	O	
ANISOU	1013	O	LEU	A	145	3258	3290	3742	304	515	436	O
ATOM	1014	CB	LEU	A	145	48.239	8.399	14.858	1.00	23.01	C	
ANISOU	1014	CB	LEU	A	145	2747	2907	3088	336	411	456	C
ATOM	1015	CG	LEU	A	145	47.852	9.914	14.837	1.00	23.44	C	
ANISOU	1015	CG	LEU	A	145	2809	2944	3152	369	450	505	C
ATOM	1016	CD1	LEU	A	145	48.449	10	627	13.574	1.00	27.23	C
ANISOU	1016	CD1	LEU	A	145	3316	3417	3614	394	503	542	C
ATOM	1017	CD2	LEU	A	145	46.301	9.907	14.788	1.00	26.07	C	
ANISOU	1017	CD2	LEU	A	145	3133	3332	3441	393	412	525	C
ATOM	1018	N	ALA	A	146	50.355	8.396	17.138	1.00	23	37	N
ANISOU	1018	N	ALA	A	146	2769	2841	3270	274	422	387	N
ATOM	1019	CA	ALA	A	146	51.005	9.050	18.186	1.00	26.88	C	
ANISOU	1019	CA	ALA	A	146	3205	3235	3775	258	436	374	C
ATOM	1020	C	ALA	A	146	52.534	8.925	17.980	1.00	26.33	C	
ANISOU	1020	C	ALA	A	146	3126	3129	3748	243	463	348	C
ATOM	1021	O	ALA	A	146	53.266	9.837	18.285	1.00	25.80	O	
ANISOU	1021	O	ALA	A	146	3054	3020	3730	236	499	350	O
ATOM	1022	CB	ALA	A	146	50.626	8.392	19.513	1.00	28.25	C	
ANISOU	1022	CB	ALA	A	146	3368	3407	3957	233	385	345	C
ATOM	1023	N	TYR	A	147	53.012	7.751	17.602	1.00	26.57	N	
ANISOU	1023	N	TYR	A	147	3151	3176	3767	235	439	317	N
ATOM	1024	CA	TYR	A	147	54.415	7.610	17.237	1.00	28.09	C	
ANISOU	1024	CA	TYR	A	147	3329	3347	3997	228	468	291	C
ATOM	1025	C	TYR	A	147	54.860	8.581	16.192	1.00	26.55	C	
ANISOU	1025	C	TYR	A	147	3139	3151	3799	242	538	325	C
ATOM	1026	O	TYR	A	147	55.881	9.262	16.394	1.00	25.72	O	

TABLE 3-continued

ANISOU 1026 O TYR A 147	3016 3008 3749 227 578 319 O
ATOM 1027 CB TYR A 147	54.765 6.129 16.804 1.00 25.81 C
ANISOU 1027 CB TYR A 147	3038 3080 3688 228 432 250 C
ATOM 1028 CG TYR A 147	56.123 6.043 16.124 1.00 25.23 C
ANISOU 1028 CG TYR A 147	2946 3000 3642 232 474 226 C
ATOM 1029 CD1 TYR A 147	57.293 5.875 16.865 1.00 26.74 C
ANISOU 1029 CD1 TYR A 147	3107 3155 3899 216 468 186 C
ATOM 1030 CD2 TYR A 147	56.218 6.213 14.708 1.00 24.58 C
ANISOU 1030 CD2 TYR A 147	2873 2951 3516 253 523 246 C
ATOM 1031 CE1 TYR A 147	58.573 5.801 16.251 1.00 29.25 C
ANISOU 1031 CE1 TYR A 147	3396 3471 4245 220 509 160 C
ATOM 1032 CE2 TYR A 147	57.491 6.158 14.099 1.00 27.09 C
ANISOU 1032 CE2 TYR A 147	3167 3268 3857 255 569 222 C
ATOM 1033 CZ TYR A 147	58.633 5.942 14.872 1.00 28.64 C
ANISOU 1033 CZ TYR A 147	3327 3431 4125 239 562 178 C
ATOM 1034 OH TYR A 147	59.841 5.930 14.301 1.00 28.36 O
ANISOU 1034 OH TYR A 147	3260 3400 4115 240 609 155 O
ATOM 1035 N ARG A 148	54.141 8.693 15.058 1.00 26.47 N
ANISOU 1035 N ARG A 148	3404 3434 3979 267 555 361 N
ATOM 1036 CA ARG A 148	54.594 9.540 14.004 1.00 27.83 C
ANISOU 1036 CA ARG A 148	3334 3354 3888 279 622 398 C
ATOM 1037 C ARG A 148	54.553 11.069 14.468 1.00 30.14 C
ANISOU 1037 C ARG A 148	3633 3598 4222 275 660 439 C
ATOM 1038 O ARG A 148	55.422 11.897 14.080 1.00 28.79 O
ANISOU 1038 O ARG A 148	3460 3396 4081 264 720 458 O
ATOM 1039 CB ARG A 148	53.705 9.376 12.728 1.00 28.21 C
ANISOU 1039 CB ARG A 148	3411 3457 3850 311 624 433 C
ATOM 1040 CG ARG A 148	53.920 8.031 12.000 1.00 28.77 C
ANISOU 1040 CG ARG A 148	3481 3572 3878 316 601 393 C
ATOM 1041 CD ARG A 148	55.376 7.936 11.483 1.00 32.31 C
ANISOU 1041 CD ARG A 148	3912 4013 4353 308 654 369 C
ATOM 1042 NE ARG A 148	55.589 8.942 10.449 1.00 29.77 N
ANISOU 1042 NE ARG A 148	3608 3700 4003 319 726 421 N
ATOM 1043 CZ ARG A 148	55.132 8.929 9.195 1.00 33.71 C
ANISOU 1043 CZ ARG A 148	4138 4248 4422 346 744 452 C
ATOM 1044 NH1 ARG A 148	54.449 7.897 8.696 1.00 29.18 N
ANISOU 1044 NH1 ARG A 148	3578 3723 3787 365 695 428 N
ATOM 1045 NH2 ARG A 148	55.368 9.994 8.428 1.00 32.89 N
ANISOU 1045 NH2 ARG A 148	4054 4142 4300 351 811 508 N
ATOM 1046 N LEU A 149	53.551 11.397 15.275 1.00 29.85 N

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TABLE 3-continued

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ANISOU	1046	N	LEU	A	149	3604	3553	4185	283	626	449	N
ATOM	1047	CA	LEU	A	149	53.488	12.756	15.962	1.00	30.95	C	
ANISOU	1047	CA	LEU	A	149	3751	3638	4371	281	651	473	C
ATOM	1048	C	LEU	A	149	54.676	13.048	16.850	1.00	31.79	C	
ANISOU	1048	C	LEU	A	149	3633	3689	4555	245	664	437	C
ATOM	1049	O	LEU	A	149	55.199	14.163	16.790	1.00	30.43	O	
ANISOU	1049	O	LEU	A	149	3667	3470	4424	235	711	460	O
ATOM	1050	CB	LEU	A	149	52.187	13.005	16.746	1.00	27.12	C	
ANISOU	1050	CB	LEU	A	149	3273	3160	3872	301	611	482	C
ATOM	1051	CG	LEU	A	149	51.002	12.987	15.708	1.00	34.00	C	
ANISOU	1051	CG	LEU	A	149	4166	4085	4669	342	605	526	C
ATOM	1052	CD1	LEU	A	149	49.597	12.944	16.307	1.00	27.23	C	
ANISOU	1052	CD1	LEU	A	149	3304	3257	3786	363	561	530	C
ATOM	1053	CD2	LEU	A	149	51.061	14.089	14.645	1.00	32.27	C	
ANISOU	1053	CD2	LEU	A	149	3977	3850	4435	367	658	585	C
ATOM	1054	N	ILE	A	150	55.060	12.091	17.686	1.00	26.86	N	
ANISOU	1054	N	ILE	A	150	3184	3059	3952	224	619	384	N
ATOM	1055	CA	ILE	A	150	56.178	12.259	18.577	1.00	28.24	C	
ANISOU	1055	CA	ILE	A	150	3333	3193	4199	192	620	345	C
ATOM	1056	C	ILE	A	150	57.480	12.411	17.783	1.00	30.56	C	
ANISOU	1056	C	ILE	A	150	3606	3482	4524	175	674	342	C
ATOM	1057	O	ILE	A	150	58.322	13.220	18.242	1.00	31.73	O	
ANISOU	1057	O	ILE	A	150	3738	3582	4737	148	701	333	O
ATOM	1058	CB	ILE	A	150	56.231	11.058	19.593	1.00	29.51	C	
ANISOU	1058	CB	ILE	A	150	3476	3371	4365	179	552	292	C
ATOM	1059	CG1	ILE	A	150	55.136	11.266	20.614	1.00	31.04	C	
ANISOU	1059	CG1	ILE	A	150	3686	3563	4544	184	515	296	C
ATOM	1060	CG2	ILE	A	150	57.644	10.918	20.292	1.00	27.11	C	
ANISOU	1060	CG2	ILE	A	150	3138	3032	4131	149	546	244	C
ATOM	1061	CD1	ILE	A	150	54.870	9.976	21.413	1.00	36.52	C	
ANISOU	1061	CD1	ILE	A	150	4375	4281	5220	173	448	261	C
ATOM	1062	N	ARG	A	151	57.587	11.759	16.601	1.00	27.66	N	
ANISOU	1062	N	ARG	A	151	3240	3159	4110	190	692	351	N
ATOM	1063	CA	ARG	A	151	58.771	11.931	15.725	1.00	29.69	C	
ANISOU	1063	CA	ARG	A	151	3476	3417	4388	176	754	351	C
ATOM	1064	C	ARG	A	151	58.931	13	394	15.284	1.00	32.49	C
ANISOU	1064	C	ARG	A	151	3848	3735	4762	165	821	405	C
ATOM	1065	O	ARG	A	151	60.070	13.921	15.230	1.00	30.84	O	
ANISOU	1065	O	ARG	A	151	3611	3497	4609	133	869	399	O
ATOM	1066	CB	ARG	A	151	58.697	11.034	14.476	1.00	27.99	C	

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TABLE 3-continued

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ANISOU 1066 CB ARG A 151	3268 3262 4106 200 764 353 C
ATOM 1067 CG ARG A 151	58.977 9.498 14.747 1.00 26.90 C
ANISOU 1067 CG ARG A 151	3107 3150 3963 206 708 290 C
ATOM 1068 CD ARG A 151	60.199 9.170 15.586 1.00 28.80 C
ANISOU 1068 CD ARG A 151	3301 3363 4280 182 694 234 C
ATOM 1069 NE ARG A 151	61.452 9.701 14.978 1.00 29.72 N
ANISOU 1069 NE ARG A 151	3381 3475 4435 165 766 232 N
ATOM 1070 CZ ARG A 151	62.049 9.233 13.878 1.00 31.50 C
ANISOU 1070 CZ ARG A 151	3592 3741 4635 177 808 222 C
ATOM 1071 NH1 ARG A 151	61.646 8.109 13.288 1.00 30.51 N
ANISOU 1071 NH1 ARG A 151	3483 3658 4450 209 778 201 N
ATOM 1072 NH2 ARG A 151	63.114 9.874 13.358 1.00 30.42 N
ANISOU 1072 NH2 ARG A 151	3421 3501 4535 155 880 228 N
ATOM 1073 N ILE A 152	57.806 14.055 14.985 1.00 25.74 N
ANISOU 1073 N ILE A 152	3543 3385 4370 191 824 458 N
ATOM 1074 CA ILE A 152	57.813 15.514 14.689 1.00 29.46 C
ANISOU 1074 CA ILE A 152	3535 3302 4355 185 879 514 C
ATOM 1075 C ILE A 152	57.734 16.461 15.867 1.00 32.97 C
ANISOU 1075 C ILE A 152	3983 3670 4855 170 356 507 C
ATOM 1076 O ILE A 152	57.645 17.690 15.631 1.00 32.84 O
ANISOU 1076 O ILE A 152	3996 3812 4868 168 905 553 O
ATOM 1077 CB ILE A 152	56.820 15.869 13.585 1.00 25.30 C
ANISOU 1077 CB ILE A 152	3563 3310 4280 225 898 580 C
ATOM 1078 CG1 ILE A 152	55.345 15.711 14.068 1.0 29.59 C
ANISOU 1078 CG1 ILE A 152	3634 3379 4267 265 838 588 C
ATOM 1079 CG2 ILE A 152	57.228 15.018 12.388 1.00 29.07 C
ANISOU 1079 CG2 ILE A 152	3542 3458 4238 231 922 580 C
ATOM 1080 CD1 ILE A 152	54.410 18.018 12.959 1.00 34.45 C
ANISOU 1080 CD1 ILE A 152	4283 4001 4835 307 856 558 C
ATOM 1081 N GLY A 153	57.844 15.924 17.368 1.00 27.89 N
ANISOU 1081 N GLY A 153	3313 3027 4255 157 813 450 N
ATOM 1082 CA GLY A 153	53.008 18.754 13.265 1.00 29.31 C
ANISOU 1082 CA GLY A 153	3491 3145 4500 136 801 429 C
ATOM 1083 C GLY A 153	56.652 17.240 18.815 1.00 33.85 C
ANISOU 1083 C GLY A 153	4103 3712 5050 172 771 447 C
ATOM 1084 O GLY A 153	56.620 13.142 19.644 1.00 82.87 O
ANISOU 1084 O GLY A 153	3988 3532 4971 184 769 433 O
ATOM 1085 N LYS A 154	55.527 16.613 18.431 1.00 30.12 N
ANISOU 1085 N LYS A 154	3644 3255 4005 211 742 466 N
ATOM 1086 CA LYS A 154	54.222 17.034 18.063 1.00 30.49 C

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TABLE 3-continued

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ANISOU 1086 CA LYS A 154	3714 3342 4527 246 714 479 C
ATOM 1087 C LYS A 154	54.011 16.287 20.271 1.30 31.69 C
ANISOU 1087 C LYS A 154	3844 3512 4686 235 650 423 C
ATOM 1088 O LYS A 154	54.481 15.074 20.403 1.00 33.10 O
ANISOU 1088 O LYS A 154	4001 3727 4861 215 627 387 O
ATOM 1089 CB LYS A 154	53.065 10.663 18.034 1.00 30.87 C
ANISOU 1089 CB LYS A 154	3781 3452 4498 290 705 521 C
ATOM 1090 CG LYS A 154	53.520 17.302 16.353 1.00 35.36 C
ANISOU 1090 CG LYS A 154	4444 4075 5101 312 755 585 C
ATOM 1091 CD LYS A 154	53.106 18.883 16.775 1.00 41.82 C
ANISOU 1091 CD LYS A 154	5230 4758 5903 318 793 621 C
ATOM 1092 CE LYS A 154	51.795 19.507 17.157 1.00 43.41 C
ANISOU 1092 CE LYS A 154	5453 4954 6067 367 703 840 C
ATOM 1093 NZ LYS A 154	51.853 20.982 10.756 1.00 35.02 N
ANISOU 1093 NZ LYS A 154	4510 3693 5130 353 313 303 N
ATOM 1094 N LYS A 155	50.230 16.806 21.181 1.00 30.54 N
ANISOU 1094 N LYS A 155	3710 3348 4545 252 637 415 N
ATOM 1095 CA LYS A 155	52.812 15.953 22.323 1.00 31.50 C
ANISOU 1095 CA LYS A 155	3814 3503 4652 244 582 370 C
ATOM 1096 C LYS A 155	51.616 15.134 21 871 1.00 31.55 C
ANISOU 1096 C LYS A 155	3820 3531 4533 271 555 391 C
ATOM 1097 O LYS A 155	50.602 15.731 21.453 1.00 31.95 O
ANISOU 1097 O LYS A 155	3887 3843 4809 310 567 423 O
ATOM 1098 CB LYS A 155	52.397 10.356 23.524 1.00 35.26 C
ANISOU 1098 CB LYS A 155	4301 3942 5154 252 574 349 C
ATOM 1099 CG LYS A 155	53.570 17.740 24.005 1.00 37.40 C
ANISOU 1099 CG LYS A 155	4574 4137 5500 221 596 324 C
ATOM 1100 CD LYS A 155	54.645 15.744 24.570 1.00 42.21 C
ANISOU 1100 CD LYS A 155	5152 4756 6130 177 534 276 C
ATOM 1101 CE LYS A 155	55.082 17.406 25.450 1.00 45.01 C
ANISOU 1101 CE LYS A 155	5499 5348 5550 142 564 233 C
ATOM 1102 NZ LYS A 155	56.283 18.597 24.753 1.00 41.30 N
ANISOU 1102 NZ LYS A 155	5103 4585 6203 133 620 280 N
ATOM 1103 N ALA A 156	51.720 13.812 21.593 1 00 33.12 N
ANISOU 1103 N ALA A 156	3622 3442 4379 252 520 369 N
ATOM 1104 CA ALA A 156	50.553 12.052 21.503 1.00 32.81 C
ANISOU 1104 CA ALA A 156	3961 3850 4857 250 490 334 C
ATOM 1105 C ALA A 156	50.455 11 856 22.512 1.00 36.84 C
ANISOU 1105 C ALA A 156	4458 4333 5158 241 439 345 C
ATOM 1106 O ALA A 156	51.504 11.229 22.920 1.00 33.79 O

TABLE 3-continued

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ANISOU	1106	O	ALA	A	156	4333	3073	4800	210	421	311	O
ATOM	1107	CB	ALA	A	156	54.775	12.384	23.110	1.00	28.91	C	
ANISOU	1107	CB	ALA	A	156	3470	3886	4129	276	501	405	C
ATOM	1108	N	ILE	A	157	49.266	11.516	22.937	1.00	38.97	N	
ANISOU	1108	N	ILE	A	157	4724	4595	5388	251	413	351	N
ATOM	1109	CA	ILE	A	157	49.018	14.433	23.757	1.00	35.83	C	
ANISOU	1109	CA	ILE	A	157	4441	4450	5393	220	364	325	C
ATOM	1110	C	ILE	A	157	47.532	9.503	23.083	1.00	40.11	C	
ANISOU	1110	C	ILE	A	157	4654	4931	5456	224	340	344	C
ATOM	1111	O	ILE	A	157	46.904	10.038	22.335	1.00	37.03	O	
ANISOU	1111	O	ILE	A	157	4457	4576	5335	254	351	372	O
ATOM	1112	CB	ILE	A	157	48.507	10.831	25.134	1.00	37.13	O	
ANISOU	1112	CB	ILE	A	157	4430	4491	5133	215	354	308	C
ATOM	1113	CG1	ILE	A	157	48.232	9.572	25.973	1.00	44.39	C	
ANISOU	1113	CG1	ILE	A	157	5390	5442	2030	173	305	288	C
ATOM	1114	CG2	ILE	A	157	47.212	11.550	24.953	1.00	45.95	C	
ANISOU	1114	CG2	ILE	A	157	5591	5342	6226	252	372	334	C
ATOM	1115	CD1	ILE	A	157	47.955	9.045	27.372	1.00	43.40	C	
ANISOU	1115	CD1	ILE	A	157	5271	5314	5905	169	298	238	C
ATOM	1116	N	MET	A	158	48.303	8.284	22.974	1.00	36.56	N	
ANISOU	1116	N	MET	A	158	4405	4493	4995	196	304	327	N
ATOM	1117	CA	MET	A	158	47.317	7.285	22.577	1.00	35.12	C	
ANISOU	1117	CA	MET	A	158	4217	4365	4752	186	270	336	C
ATOM	1118	C	MET	A	158	46.388	5.612	23.840	1.00	37.48	C	
ANISOU	1118	C	MET	A	158	4510	4684	5046	151	232	323	C
ATOM	1119	O	MET	A	158	47.405	6.107	24.701	1.00	36.58	O	
ANISOU	1119	O	MET	A	158	4406	4538	4953	122	209	298	O
ATOM	1120	CB	MET	A	158	47.982	5.317	21.614	1.00	36.83	C	
ANISOU	1120	CB	MET	A	158	4443	4579	4972	178	254	325	C
ATOM	1121	CG	MET	A	158	47.167	5.126	21.141	1.00	29.56	C	
ANISOU	1121	CG	MET	A	158	3522	3703	4005	162	212	325	C
ATOM	1122	SD	MET	A	158	45.632	5.631	20.442	1.00	34.46	S	
ANISOU	1122	SD	MET	A	158	4127	4396	4578	190	219	356	S
ATOM	1123	CE	MET	A	158	44.739	4.056	29.461	1.00	34.30	C	
ANISOU	1123	CE	MET	A	158	4102	4414	4518	145	157	352	C
ATOM	1124	N	PHE	A	159	45.363	6.600	23.953	1.00	29.65	N	
ANISOU	1124	N	PHE	A	159	3539	3784	4055	153	224	341	N
ATOM	1125	CA	PHE	A	159	44.585	5.863	25.358	1.00	35.28	C	
ANISOU	1125	CA	PHE	A	159	4208	4488	4710	112	192	334	C
ATOM	1126	C	PHE	A	159	44.052	4.531	24.476	1.00	35.95	C	

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ANISOU	1126	C	PHE	A	159	4418	4740	4387	81	151	340	C
ATOM	1127	O	PHE	A	159	43.182	4.653	23.552	1.00	44.07	0	
ANISOU	1127	O	PHE	A	159	5299	5536	5755	100	154	359	C
ATOM	1128	CB	PHE	A	159	43.541	6.723	25.597	1.00	32.15	15	C
ANISOU	1128	CB	PHE	A	159	3824	4174	4332	132	216	343	C
ATOM	1129	CG	PHE	A	159	43.885	8.064	26.185	1.00	27.80	15	C
ANISOU	1129	CG	PHE	A	159	3239	3550	3774	166	254	339	C
ATOM	1130	CD1	PHE	A	159	44.005	9.104	25.377	1.00	30.52	15	C
ANISOU	1130	CD1	PHE	A	159	3584	3874	4137	215	291	354	C
ATOM	1131	CD2	PHE	A	159	44.050	8.209	27.570	1.00	34.76	15	C
ANISOU	1131	CD2	PHE	A	159	4126	4413	4653	146	253	317	C
ATOM	1132	CE1	PHE	A	159	44.276	10.457	25	934	1.00	32.09	C
ANISOU	1132	CE1	PHE	A	159	3790	4034	4368	245	324	345	C
ATOM	1133	CE2	PHE	A	159	44.381	9.442	28.104	1.00	33.45	15	C
ANISOU	1133	CE2	PHE	A	159	3665	4217	4526	176	285	303	C
ATOM	1134	CZ	PHE	A	159	44.465	10.554	27.290	1.00	36.63	15	C
ANISOU	1134	CZ	PHE	A	159	4359	4505	4952	225	320	317	C
ATOM	1135	N	GLU	A	160	44.320	3.373	24.698	1.00	47.87	16	N
ANISOU	1135	N	GLU	A	160	5816	6138	6266	34	110	326	N
ATOM	1136	CA	GLU	A	160	48.175	2.339	24.855	1.00	43.24	16	C
ANISOU	1136	CA	GLU	A	160	5217	5572	5639	-5	74	338	C
ATOM	1137	C	GLU	A	160	42.372	2.121	25.151	1.00	51.40	16	C
ANISOU	1137	C	GLU	A	160	6239	6637	6652	-46	66	346	C
ATOM	1138	O	GLU	A	160	41.263	1.315	26.125	1.00	42.89	16	O
ANISOU	1138	O	GLU	A	160	5144	5609	5542	-88	40	659	O
ATOM	1139	CB	GLU	A	160	43.592	0.941	24.284	1.00	47.84	16	C
ANISOU	1139	CB	GLU	A	160	5526	5122	6219	-37	25	225	C
ATOM	1140	CG	GLU	A	160	44.437	1.075	23.047	1.00	41.36	16	C
ANISOU	1140	CG	GLU	A	160	5318	5285	5113	2	37	312	C
ATOM	1141	CD	GLU	A	160	45.857	1.410	23.356	0.50	40.71	16	C
ANISOU	1141	CD	GLU	A	160	4953	5141	5373	20	53	292	C
ATOM	1142	OE1	GLU	A	160	46.200	1.513	24.566	1.00	43.83	16	O
ANISOU	1142	OE1	GLU	A	160	5355	5512	5788	2	48	287	O
ATOM	1143	OE2	GLU	A	160	46.336	1.539	22.411	1.00	38.00	16	O
ANISOU	1143	OE2	GLU	A	160	4615	4773	5343	40	58	282	O
ATOM	1144	N	ASP	A	161	42.879	2.756	27.262	1.00	36.99	16	N
ANISOU	1144	N	ASP	A	161	4423	4786	4845	-41	85	337	N
ATOM	1145	CA	ASP	A	161	41.975	2.858	28.440	1.00	30.48	16	C
ANISOU	1145	CA	ASP	A	161	3582	4006	3993	-88	93	345	C
ATOM	1146	C	ASP	A	161	40.900	3.942	28.171	1.00	29.48	16	C

TABLE 3-continued

ANISOU 1146	CASP A	161	3411	3938	3852	-26	134	357	C
ATOM	1147	O ASP A	161	41.221	5.144	28.240	1.00	28.77	O
ANISOU	1147	O ASP A	161	3318	3829	3783	24	170	349
ATOM	1148	CB ASP A	161	42.696	3.087	29.769	1.00	28.81	C
ANISOU	1148	CB ASP A	161	3396	3759	-79	95	329	C
ATOM	1149	CG ASP A	161	41.722	2.969	30.962	1.00	29.50	C
ANISOU	1149	CGASP A	161	3470	3901	3839	-115	101	339
ATOM	1150	OD1 ASP A	161	42.057	2.584	32.173	1.00	30.23	O
ANISOU	1150	OD1 ASP A	161	3590	3978	3917	-151	85	335
ATOM	1151	OD2 ASP A	161	40.504	3.255	30.684	1.00	31.66	O
ANISOU	1151	OD2 ASP A	161	3700	4240	4088	-108	123	353
ATOM	1152	N THR A	162	39.695	3.538	27.876	1.00	29.79	N
ANISOU	1152	N THR A	162	3415	4043	3860	-44	126	373
ATOM	1153	CA THR A	162	38.622	4.475	27.394	1.00	30.80	C
ANISOU	1153	CA THR A	162	3494	4232	3976	3	157	384
ATOM	1154	C THR A	162	38.028	5.248	28.556	1.00	33.29	C
ANISOU	1154	C THR A	162	3785	4582	4282	15	192	379
ATOM	1155	O THR A	162	37.404	6.280	28.339	1.00	31.23	O
ANISOU	1155	O THR A	162	3492	4353	4022	70	223	381
ATOM	1156	CB THR A	162	37.551	3.755	26.588	1.00	33.20	C
ANISOU	1156	CB THR A	162	3763	4599	4254	-19	131	400
ATOM	1157	OG1 THR A	162	37.219	2.576	27.292	1.00	34.72	O
ANISOU	1157	OG1 THR A	162	3957	4809	4426	-96	103	403
ATOM	1158	CG2 THR A	162	38.119	3.279	25.205	1.00	30.39	C
ANISOU	1158	CG2 THR A	162	3431	4211	3905	-9	104	400
ATOM	1159	N HIS A	163	38.271	4.780	29.711	1.00	30.54	N
ANISOU	1159	N HIS A	163	3456	4225	3922	-32	186	370
ATOM	1160	CA HIS A	163	37.975	5.534	31.010	1.00	27.07	C
ANISOU	1160	CA HIS A	163	3006	3808	3470	-20	221	357
ATOM	1161	C HIS A	163	38.869	6.703	31.137	1.00	29.09	C
ANISOU	1161	C HIS A	163	3287	4004	3761	36	246	336
ATOM	1162	O HIS A	163	38.382	7.842	31.299	1.00	28.18	O
ANISOU	1162	O HIS A	163	3149	3909	3649	89	284	326
ATOM	1163	CB HIS A	163	38.074	4.610	32.212	1.00	30.05	C
ANISOU	1163	CBHIS A	163	3407	4190	3819	-90	203	358
ATOM	1164	CG HIS A	163	36.906	3.665	32.281	-149	1.00	31.76
ANISOU	1164	CG HIS A	163	3589	4481	3999	-149	192	381
ATOM	1165	ND1 HIS A	163	36.022	3.671	33.312	1.00	30.75	N
ANISOU	1165	ND1HIS A	163	3432	4421	3832	-177	217	384
ATOM	1166	CD2 HIS A	163	36.483	2.678	31.377	1.00	29.90	C

TABLE 3-continued

ANISOU	1166	CD2HIS	A	163	3339	4261	3759	-187	157	400	C	
ATOM	1167	CE1	HIS	A	163	35.081	2.686	33.110	1.00	29.33	C	
ANISOU	1167	CE1	HIS	A	163	3220	4298	3627	-238	200	408	C
ATOM	1168	NE2	HIS	A	163	35.352	2.103	31.892	1.00	30.65	N	
ANISOU	1168	NE2	HIS	A	163	3396	4431	3818	-242	161	416	N
ATOM	1169	N	LEU	A	164	40.184	6.417	30.952	1.00	28.37	N	
ANISOU	1169	N	LEU	A	164	3243	3837	3701	28	225	327	N
ATOM	1170	CA	LEU	A	164	41.153	7.566	31.008	1.00	32.27	C	
ANISOU	1170	CA	LEU	A	164	3759	4267	4235	74	247	306	C
ATOM	1171	C	LEU	A	164	40.902	8.537	2.848	1.00	29.85	C	
ANISOU	1171	CLEU	A	164	3436	3959	3952	136	274	318	C	
ATOM	1172	O	LEU	A	164	40.979	9.789	30.036	1.00	25.42	O	
ANISOU	1172	O	LEU	A	164	2875	3374	3411	186	308	306	O
ATOM	1173	CB	LEU	A	164	42.577	7.044	30.949	1.00	31.08	C	
ANISOU	1173	CB	LEU	A	164	3650	4044	4114	51	218	295	C
ATOM	1174	CG	LEU	A	164	43.134	6.444	32.261	1.00	40.89	C	
ANISOU	1174	CG	LEU	A	164	4922	5268	5345	6	192	278	C
ATOM	1175	CD1	LEU	A	164	44.021	5.082	32.070	1.00	40.50	C	
ANISOU	1175	CD1	LEU	A	164	4906	5145	5337	-2	163	263	C
ATOM	1176	CD2	LEU	A	164	43.022	7.431	33.413	1.00	40.21	C	
ANISOU	1176	CD2	LEU	A	164	4838	5188	5253	25	221	254	C
ATOM	1177	N	ALA	A	165	40.628	7.560	28.652	1.00	27.50	N	
ANISOU	1177	N	ALA	A	165	3138	3691	3653	134	257	340	N
ATOM	1178	CA	ALA	A	165	40.430	6.812	27.465	1.00	29.72	C	
ANISOU	1178	CA	ALA	A	165	3396	3957	3940	193	278	358	C
ATOM	1179	C	ALA	A	165	39.182	9.717	27.739	1.00	29.80	C	
ANISOU	1179	C	ALA	A	165	3366	4023	3035	239	306	352	C
ATOM	1180	O	ALA	A	165	39.102	10.942	27.441	1.00	27.67	O	
ANISOU	1180	O	ALA	A	165	3098	3727	3687	333	335	354	O
ATOM	1181	CB	ALA	A	165	40.138	7.925	26.245	1.00	27.53	C	
ANISOU	1181	CB	ALA	A	165	3110	3708	3544	183	249	379	C
ATOM	1182	N	ALA	A	166	38.115	9.115	28.282	1.00	29.86	N	
ANISOU	1182	N	ALA	A	166	3335	4104	3935	209	296	363	N
ATOM	1183	CA	ALA	A	166	36.538	9.958	28.698	1.00	29.65	C	
ANISOU	1183	CA	ALA	A	166	3252	4140	3565	254	326	360	C
ATOM	1184	C	ALA	A	166	37.299	11.060	29.686	1.00	27.62	C	
ANISOU	1184	C	ALA	A	166	3324	3842	3626	289	363	331	C
ATOM	1185	O	ALA	A	166	36.531	12.217	29.530	1.00	28.30	O	
ANISOU	1185	O	ALA	A	166	3095	3930	3727	357	348	328	O
ATOM	1186	CB	ALA	A	166	35.755	9.150	29.257	1.0	25.24	C	

TABLE 3-continued

ANISOU	1186	CB	ALA A	166	2655	3671	3265	209	318	362	C
ATOM	1187	N	MET A	167	38.055	10.723	30.747	1.00	33.10	N	
ANISOU	1187	N	MET A	167	3383	4135	3954	244	356	309	N
ATOM	1188	CA	MET A	167	38.437	11.733	31.712	1.00	28.03	C	
ANISOU	1188	CA	MET A	167	3130	3825	3597	274	384	276	C
ATOM	1189	C	MET A	167	39.268	12.861	21.112	100	28.40	C	
ANISOU	1189	C	MET A	167	3208	3789	3795	326	400	272	C
ATOM	1190	O	MET A	167	38.036	14.362	31.347	1.00	32.41	O	
ANISOU	1190	O	MET A	167	3711	4284	4318	385	430	255	O
ATOM	1191	CB	MET A	167	39.103	11.107	32.969	1.00	28.43	C	
ANISOU	1191	CB	MET A	167	3211	3860	3731	215	370	252	C
ATOM	1192	CG	MET A	167	38.145	10.379	33.517	1.00	48.40	C	
ANISOU	1192	CG	MET A	167	3176	3942	3571	161	352	263	C
ATOM	1193	SD	MET A	167	30.140	9.503	35.007	1.00	31.56	S	
ANISOU	1193	SD	MET A	167	3529	4307	4056	103	341	243	S
ATOM	1194	CE	MET A	167	37.838	8.657	35.917	1.00	32.72	C	
ANISOU	1194	CE	MET A	167	3740	4559	4132	50	349	253	C
ATOM	1195	N	SER A	168	40.256	12.529	30.271	1.00	26.98	N	
ANISOU	1195	N	SER A	168	3056	3553	3541	339	382	287	N
ATOM	1196	CA	SER A	168	41.233	13.603	29.570	1.60	29.45	C	
ANISOU	1196	CA	SER A	168	3398	3789	4034	353	401	291	C
ATOM	1197	C	SER A	168	40.222	14.452	28.580	1.30	28.04	C	
ANISOU	1197	C	SER A	168	3201	3626	38241	420	421	317	C
ATOM	1198	O	SER A	168	40.320	15.685	28.660	1.00	29.18	O	
ANISOU	1198	O	SER A	168	3360	3723	4303	471	443	311	O
ATOM	1199	CB	SER A	168	42.240	13.044	28.741	1.00	29.97	C	
ANISOU	1199	CB	SER A	168	349	3804	4094	322	384	304	C
ATOM	1200	OG	SER A	168	43.272	12.925	29.663	1.00	48.80	O	
ANISOU	1200	OG	SER A	168	5900	6141	5499	287	375	272	O
ATOM	1201	N	ALA A	169	39.388	13.817	27.954	1.00	28.22	N	
ANISOU	1201	N	ALA A	169	3191	3712	3813	421	405	345	N
ATOM	1202	CA	ALA A	169	38.537	14.569	26.595	1.00	29.82	C	
ANISOU	1202	CA	ALA A	169	3373	3938	4018	489	416	375	C
ATOM	1203	CA	ALA A	169	37.607	15.491	27.659	1.00	32.72	C	
ANISOU	1203	CA	ALA A	169	3715	4333	4385	542	440	355	C
ATOM	1204	O	ALA A	169	37.311	16.603	27.205	1.00	31.83	O	
ANISOU	1204	O	ALA A	169	3607	4195	4293	612	457	367	O
ATOM	1205	CB	ALA A	169	37.705	13.615	26.011	1.00	30.28	C	
ANISOU	1205	CB	ALA A	169	3395	4073	4037	475	387	403	C
ATOM	1206	N	SER A	170	37.151	15.054	28.842	1.00	28.98	N	

TABLE 3-continued

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ANISOU	1206	N	SER	A	170	3215	3909	3886	511	441	325	N
ATOM	1207	CA	SER	A	170	36.099	15.838	29.498	1.00	34.32	C	
ANISOU	1207	CA	SER	A	170	3856	4630	4555	565	466	304	C
ATOM	1208	C	SER	A	170	36.670	17.168	30.044	1.00	31.85	C	
ANISOU	1208	C	SER	A	170	3584	4235	4283	614	495	274	C
ATOM	1209	O	SER	A	170	35.907	18.059	30.306	1.00	31.66	O	
ANISOU	1209	O	SER	A	170	3540	4227	4264	679	516	259	O
ATOM	1210	CB	SER	A	170	35.387	15.098	30.658	1.00	31.43	C	
ANISOU	1210	CB	SER	A	170	3449	4347	4145	521	469	279	C
ATOM	1211	OG	SER	A	170	36.323	15.072	31.677	1.00	30.11	O	
ANISOU	1211	OG	SER	A	170	3325	4129	3985	483	474	248	O
ATOM	1212	N	ARG	A	171	37.991	17.288	30.132	1.00	32.38	N	
ANISOU	1212	N	ARG	A	171	3705	4214	4383	583	493	266	N
ATOM	1213	CA	ARG	A	171	38.603	18.496	30.673	1.00	32.42	C	
ANISOU	1213	CA	ARG	A	171	3751	4137	4431	618	515	235	C
ATOM	1214	C	ARG	A	171	39.411	19.189	29.569	1.00	34.25	C	
ANISOU	1214	C	ARG	A	171	4024	4280	4711	640	520	266	C
ATOM	1215	O	ARG	A	171	40.203	20.099	29.789	1.00	34.65	O	
ANISOU	1215	O	ARG	A	171	4116	4242	4806	652	535	248	O
ATOM	1216	CB	ARG	A	171	39.494	18.175	31.881	1.00	33.97	C	
ANISOU	1216	CB	ARG	A	171	3974	4305	4628	560	510	191	C
ATOM	1217	CG	ARG	A	171	38.746	17.520	33.085	1.00	38.76	C	
ANISOU	1217	CG	ARG	A	171	4549	4998	5180	533	510	162	C
ATOM	1218	CD	ARG	A	171	38.162	18.596	34.069	1.00	45.95	C	
ANISOU	1218	CD	ARG	A	171	5455	5914	6089	590	541	113	C
ATOM	1219	NE	ARG	A	171	37.255	19.510	33.465	1.00	55.37	N	
ANISOU	1219	NE	ARG	A	171	6625	7116	7297	672	561	124	N
ATOM	1220	CZ	ARG	A	171	36.987	20.776	33.832	1.00	63.43	C	
ANISOU	1220	CZ	ARG	A	171	7657	8099	8343	744	586	90	C
ATOM	1221	NH1	ARG	A	171	37.601	21.417	34.823	1.00	63.82	N	
ANISOU	1221	NH1	ARG	A	171	7747	8092	8409	745	597	37	N
ATOM	1222	NH2	ARG	A	171	36.081	21.421	33.140	1.00	65.34	N	
ANISOU	1222	NH2	ARG	A	171	7872	8359	8595	820	597	109	N
ATOM	1223	N	SER	A	172	39.221	18.735	28.369	1.00	31.33	N	
ANISOU	1223	N	SER	A	172	3643	3934	4328	642	507	314	N
ATOM	1224	CA	SER	A	172	39.953	19.218	27.240	1.00	31.69	C	
ANISOU	1224	CA	SER	A	172	3725	3909	4405	656	513	351	C
ATOM	1225	C	SER	A	172	39.697	20.738	26.972	1.00	35.30	C	
ANISOU	1225	C	SER	A	172	4208	4306	4900	735	537	360	C
ATOM	1226	O	SER	A	172	38.559	21.235	27.208	1.00	33.82	O	

TABLE 3-continued

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ANISOU	1226	O	SER	A	172	3992	4159	4700	797	542	353	O
ATOM	1227	CB	SER	A	172	39.455	18.464	26.052	1.00	32.63	C	
ANISOU	1227	CB	SER	A	172	3821	4088	4489	655	493	396	C
ATOM	1228	OG	SER	A	172	40.415	18.526	25.004	1.00	36.27	O	
ANISOU	1228	OG	SER	A	172	4320	4492	4970	643	497	430	O
ATOM	1229	N	SER	A	173	40.738	21.446	26.529	1.00	33.86	N	
ANISOU	1229	N	SER	A	173	4076	4026	4765	732	553	375	N
ATOM	1230	CA	SER	A	173	40.688	22.937	26.506	1.00	36.21	C	
ANISOU	1230	CA	SER	A	173	4409	4242	5106	796	576	377	C
ATOM	1231	C	SER	A	173	41.407	23.425	25.263	1.00	37.70	C	
ANISOU	1231	C	SER	A	173	4642	4361	5323	799	583	432	C
ATOM	1232	O	SER	A	173	42.032	22.641	24.543	1.00	34.38	O	
ANISOU	1232	O	SER	A	173	4221	3953	4887	750	583	459	O
ATOM	1233	CB	SER	A	173	41.382	23	525	27.734	1.00	39.74	C
ANISOU	1233	CB	SER	A	173	4882	4321	5595	775	588	317	C
ATOM	1234	OG	SER	A	173	42.704	23.277	27.585	1.00	36.10	O	
ANISOU	1234	OG	SER	A	173	4461	4112	5177	706	590	314	O
ATOM	1235	N	GLN	A	174	41.225	24.719	24.973	1.00	37.88	N	
ANISOU	1235	N	GLN	A	174	4700	4311	5381	352	605	451	N
ATOM	1236	CA	GLN	A	174	41.853	25.330	23.306	1.00	38.15	C	
ANISOU	1236	CA	GLN	A	174	4735	4270	5442	869	623	511	C
ATOM	1237	C	GLN	A	174	43.295	24.935	23.613	1.00	32.75	C	
ANISOU	1237	C	GLN	A	174	4121	3540	4782	766	636	513	C
ATOM	1238	O	GLN	A	174	44.071	25.049	24.541	1.00	39.85	O	
ANISOU	1238	O	GLN	A	174	5001	4371	5592	741	541	454	O
ATOM	1239	CB	GLN	A	174	41.788	23.930	23.953	1.00	40.04	C	
ANISOU	1239	CB	GLN	A	174	5112	4443	5774	930	642	514	C
ATOM	1240	CG	GLN	A	174	40.491	27.497	23.424	1.00	53.58	C	
ANISOU	1240	CG	GLN	A	174	6795	5150	7442	1023	631	543	C
ATOM	1241	CD	GLN	A	174	40.235	27.331	22.022	1.00	51.34	C	
ANISOU	1241	CD	GLN	A	174	6494	5913	7398	1037	620	620	C
ATOM	1242	OE1	GLN	A	174	39.128	26.545	21.719	1.00	57.40	O	
ANISOU	1242	OE1	GLN	A	174	7218	6774	7818	1082	595	632	O
ATOM	1243	NE2	GLN	A	174	41.286	26.950	21.191	1.00	49.80	N	
ANISOU	1243	NE2	GLN	A	174	6277	5504	5851	988	538	554	N
ATOM	1244	N	GLY	A	175	43.623	24.535	22.400	1.00	33.94	N	
ANISOU	1244	N	GLY	A	175	4280	3705	4905	770	640	568	N
ATOM	1245	CA	GLY	A	175	44.548	24.1142	22.031	1.00	33.72	C	
ANISOU	1245	CA	GLY	A	175	4290	3672	4925	697	656	573	C
ATOM	1246	C	GLY	A	175	45.007	22.594	21.957	1.00	33	10	C

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TABLE 3-continued

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ANISOU	1246	C	GLY	A	175	4142	3661	4772	630	631	559	C
ATOM	1247	O	GLY	A	175	45.972	22.021	21.445	1.00	35.80	O	
ANISOU	1247	O	GLY	A	175	4486	3939	5116	598	640	558	O
ATOM	1248	N	ASP	A	176	43.995	21.908	22.493	1.00	31.91	N	
ANISOU	1248	N	ASP	A	176	3950	3593	4580	567	501	584	N
ATOM	1249	CA	ASP	A	176	44.000	20.391	22.341	1.00	31.95	C	
ANISOU	1249	CA	ASP	A	176	3919	3652	4539	621	574	525	C
ATOM	1250	C	ASP	A	176	43.399	20.025	25.956	1.00	33.53	C	
ANISOU	1250	C	ASP	A	176	3736	3557	4308	550	565	579	C
ATOM	1251	O	ASP	A	176	42.579	20.767	20.498	1.00	29.53	O	
ANISOU	1251	O	ASP	A	176	3618	3433	4169	713	567	615	O
ATOM	1252	CB	ASP	A	176	43.123	19.701	23.420	1.00	32.11	C	
ANISOU	1252	CB	ASP	A	176	3307	3773	4531	617	545	481	C
ATOM	1253	CG	ASP	A	176	43.529	19.940	24.801	1.00	33.22	C	
ANISOU	1253	CG	ASP	A	176	4053	3887	4720	589	549	423	C
ATOM	1254	OD1	ASP	A	176	44.835	20.205	24.885	1.00	34.73	O	
ANISOU	1254	OD1	ASP	A	176	4280	3791	4945	555	585	416	O
ATOM	1255	OD2	ASP	A	176	42.873	19.722	25.811	1.50	33.56	O	
ANISOU	1255	OD2	ASP	A	176	4057	3964	4730	595	535	390	O
ATOM	1256	N	LEU	A	177	43.579	13.819	20.454	1.00	31.244	N	
ANISOU	1256	N	LEU	A	177	3839	3697	4262	608	547	580	N
ATOM	1257	CA	LEU	A	177	43.038	18.256	19.263	1.00	30.24	C	
ANISOU	1257	CA	LEU	A	177	3675	3537	4176	629	529	619	C
ATOM	1258	C	LEU	A	177	42.925	13.724	19.538	1.00	29.01	C	
ANISOU	1258	C	LEU	A	177	3463	3552	3989	579	490	585	C
ATOM	1259	O	LEU	A	177	43.842	16.138	20.109	1.00	29.60	O	
ANISOU	1259	O	LEU	A	177	3555	5604	4085	523	492	653	O
ATOM	1260	CB	LEU	A	177	43.984	18.392	18.052	1.00	29.50	C	
ANISOU	1260	CB	LEU	A	177	3813	3503	4082	613	556	853	C
ATOM	1261	CG	LEU	A	177	43.752	17.697	16.733	1.00	33.30	C	
ANISOU	1261	CG	LEU	A	177	4399	4354	4500	622	540	690	C
ATOM	1262	CD1	LEU	A	177	42.512	13.214	15.104	1.00	35.45	C	
ANISOU	1262	CD1	LEU	A	177	4374	4351	4735	694	525	734	C
ATOM	1263	CD2	LEU	A	177	44.857	17.804	15.730	1.00	37.01	C	
ANISOU	1263	CD2	LEU	A	177	4503	4484	4970	506	678	723	C
ATOM	1264	N	TRP	A	178	41.306	16.135	19.142	1.00	25.45	N	
ANISOU	1264	N	TRP	A	178	3129	3310	3615	597	460	595	N
ATOM	1265	CA	TRP	A	178	41.591	14.544	19.343	1.00	26.50	C	
ANISOU	1265	CA	TRP	A	178	3115	3391	3800	546	422	551	C
ATOM	1266	C	TRP	A	178	41.455	13.905	13.352	1.30	29.07	C	

TABLE 3-continued

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ANISOU	1266	C	TRP	A	178	3432	3750	3865	545	402	590	C
ATOM	1267	O	TRP	A	178	40.793	14.785	17.147	1.00	33.38	O	
ANISOU	1267	O	TRP	A	178	4019	4360	4419	595	399	623	O
ATOM	1268	CB	TRP	A	178	40.327	14.387	20.111	1.00	35.77	C	
ANISOU	1268	CB	TRP	A	178	3092	3478	3503	555	357	553	C
ATOM	1269	CG	TRP	A	178	40.540	14.375	21.497	1.00	29.00	C	
ANISOU	1269	CG	TRP	A	178	3772	3722	3925	545	414	516	C
ATOM	1270	CD1	TRP	A	178	43.393	16.154	21.995	1.00	31.35	C	
ANISOU	1270	CD1	TRP	A	178	3341	3977	4256	593	443	515	C
ATOM	1271	CD2	TRP	A	178	41.021	14.097	22.554	1.00	27.71	C	
ANISOU	1271	CD2	TRP	A	178	3202	3554	3771	485	404	475	C
ATOM	1272	NE1	TAP	A	178	40.755	16.232	23.214	1.00	32.60	N	
ANISOU	1272	NE1	TRP	A	178	3833	4109	4441	5814	455	473	N
ATOM	1273	CE2	TRP	A	178	41.143	14.999	27.723	1.00	30.22	C	
ANISOU	1273	CE2	TRP	A	178	3525	2331	4125	495	427	450	C
ATOM	1274	CE3	TRP	A	178	41.303	12.718	22.717	1.80	29.02	C	
ANISOU	1274	CE3	TRP	A	178	7360	7747	7918	424	373	453	C
ATOM	1275	CZ2	TRP	A	178	41.545	14.535	24.092	1.00	29.73	C	
ANISOU	1275	CZ2	TRP	A	178	3462	3761	4073	4.52	420	409	C
ATOM	1276	CZ3	TRP	A	178	41.729	12.269	23.977	1.00	27.33	C	
ANISOU	1276	CZ3	TRP	A	178	3153	3525	3728	313	356	421	C
ATOM	1277	CH2	TRP	A	178	41.819	13.162	25	595	1.30	29.10	C
ANISOU	1277	CH2	TRP	A	178	3375	3710	3971	392	339	393	C
ATOM	1278	N	PHE	A	179	42.055	12.709	17.972	1.00	23.31	N	
ANISOU	1278	N	PHE	A	179	3835	3654	3163	492	383	555	N
ATOM	1279	CA	PHE	A	179	41.732	11.798	16.380	1.00	27.50	C	
ANISOU	1279	CA	PHE	A	179	3230	3614	3604	436	355	514	C
ATOM	1280	C	PHE	A	179	41.012	10.572	17.408	1.00	31.54	C	
ANISOU	1280	C	PHE	A	179	3742	4224	4130	445	307	544	C
ATOM	1281	O	PHE	A	179	41.451	9.958	18.419	1.00	29.54	O	
ANISOU	1281	O	PHE	A	179	3453	3953	3377	397	298	611	O
ATOM	1282	CB	PHE	A	179	43.103	11.316	18.357	1.00	20.14	C	
ANISOU	1282	CB	PHE	A	179	3594	3910	3948	455	370	554	C
ATOM	1283	CG	PHE	A	179	43.723	12.295	15.390	1.00	28.88	C	
ANISOU	1283	CG	PHE	A	179	3471	3714	3738	440	415	503	C
ATOM	1284	CD1	PHE	A	179	44.549	13.293	15.872	1.00	20.44	C	
ANISOU	1284	CD1	PHE	A	179	3433	3595	3797	492	459	609	C
ATOM	1285	CD2	PHE	A	179	43.460	12.241	14.029	1.00	29.14	C	
ANISOU	1285	CD2	PHE	A	179	3520	3734	3755	519	412	635	C
ATOM	1286	CE1	PHE	A	179	45.156	14.202	15.020	1.00	23.30	C	

TABLE 3-continued

ANISOU	1286	CE1	PHE	A	179	3451	3528	3775	515	503	648	C	
ATOM	1287	CE2	PHE	A	179	44.057	13.174	13.175	1.00	28.55	C		
ANISOU	1287	CE2	PHE	A	179	9563	3673	3590	541	457	678	C	
ATOM	1288	CZ	PHE	A	179	44.335	14.176	13.574	1.55	29.33	C		
ANISOU	1288	CZ	PHE	A	179	3500	3597	3346	545	503	688	C	
ATOM	1289	N	ALA	A	180	39.909	10.243	15.771	1.00	29.60	N		
ANISOU	1289	N	ALA	A	180	3460	4335	3326	452	274	558	N	
ATOM	1290	CA	ALA	A	180	39.069	9.039	17.178	1.00	28.617	C		
ANISOU	1290	CA	ALA	A	180	3278	3953	3561	419	226	534	C	
ATOM	1291	C	ALA	A	180	39.349	8.097	15.075	1.00	29.85	C		
ANISOU	1291	C	ALA	A	180	3445	4121	3773	398	1	99	528	C
ATOM	1292	O	ALA	A	180	33.935	8.355	14.942	1.00	29.17	O		
ANISOU	1292	O	ALA	A	180	3366	4066	3650	438	192	553	O	
ATOM	1293	CB	ALA	A	180	37.535	9.531	17.207	1.00	24.90	C		
ANISOU	1293	CB	ALA	A	180	2752	3549	3151	459	210	550	C	
ATOM	1294	N	VAL	A	181	40.025	5.939	15.400	1.00	233.63	N		
ANISOU	1294	N	VAL	A	181	3305	3943	3629	342	182	494	N	
ATOM	1295	CA	VAL	A	181	46.398	5.943	15.429	1.00	27.68	C		
ANISOU	1295	CA	VAL	A	181	3205	3836	3478	321	153	476	C	
ATOM	1296	C	VAL	A	181	39.442	4.748	19.591	1.00	23.93	C		
ANISOU	1296	C	VAL	A	181	3336	4045	3512	275	97	458	C	
ATOM	1297	O	VAL	A	181	39.484	4.052	15.611	1.30	26.59	O		
ANISOU	1297	O	VAL	A	181	3343	3747	3351	223	80	438	O	
ATOM	1298	CB	VAL	A	181	41.334	5.440	15.589	1.00	30.79	C		
ANISOU	1298	CB	VAL	A	181	3604	4163	3802	293	168	478	C	
ATOM	1299	CG1	VAL	A	181	42.207	4.457	14.425	1.00	26.28	C		
ANISOU	1299	CG1	VAL	A	181	3087	3504	3294	285	144	431	C	
ATOM	1300	CG2	VAL	A	181	42.755	6.695	15.551	1.50	33.57	C		
ANISOU	1300	CG2	VAL	A	181	4005	4450	4289	328	228	470	C	
ATOM	1301	N	SER	A	182	38.520	4.500	14.593	1.00	27.13	N		
ANISOU	1301	N	SER	A	182	3095	3877	3337	289	55	455	N	
ATOM	1302	CA	SER	A	182	37.554	3.436	14.668	1.90	23.14	C		
ANISOU	1302	CA	SER	A	182	3189	4062	3442	244	9	448	C	
ATOM	1303	C	SER	A	182	37.151	3.171	13.210	1.00	28.95	C		
ANISOU	1303	C	SER	A	182	3298	4212	3483	267	-25	449	C	
ATOM	1304	O	SER	A	182	35.639	4.087	12.549	1.05	28.83	O		
ANISOU	1304	O	SER	A	182	3274	4231	3448	325	-15	481	O	
ATOM	1305	CB	SER	A	182	35.355	4.040	15.423	1.00	29.12	C		
ANISOU	1305	CB	SER	A	182	3255	4238	3572	251	9	467	C	
ATOM	1306	OG	SER	A	182	35.273	3.111	15.454	1.00	26.35	O		

TABLE 3-continued

ANISOU	1306	OG	SER	A	182	3076	4163	3415	204	-41	453	O
ATOM	1307	N	SER	A	183	37.263	1.932	12.750	1.30	25.98	N	
ANISOU	1307	N	SER	A	183	2539	3839	3093	224	-58	416	N
ATOM	1308	CA	SER	A	183	36.850	1.603	11.355	1.65	29.34	C	
ANISOU	1308	CA	SER	A	183	3374	4315	3453	244	-105	410	C
ATOM	1309	C	SER	A	183	35.349	1.777	11.254	1.00	32.69	C	
ANISOU	1309	C	SER	A	183	3704	4340	3885	249	-143	425	C
ATOM	1310	O	SER	A	183	34.852	2.254	10.243	1.00	32.24	O	
ANISOU	1310	O	SER	A	183	3573	4815	3757	208	-155	444	O
ATOM	1311	CB	SER	A	183	37.157	0.127	11.053	1.00	29.54	C	
ANISOU	1311	CB	SER	A	183	3425	4324	3474	189	-151	361	C
ATOM	1312	OG	SER	A	183	38.518	0.000	10.626	1.00	31.35	O	
ANISOU	1312	OG	SER	A	183	3703	4495	3707	205	-122	344	O
ATOM	1313	N	SER	A	184	34.529	1.340	12.310	1.30	31.44	N	
ANISOU	1313	N	SER	A	184	3536	4581	3730	198	-161	418	N
ATOM	1314	CA	SER	A	184	33.159	1.369	12.265	1.00	30.85	C	
ANISOU	1314	CA	SER	A	184	3395	4690	3638	191	-198	426	C
ATOM	1315	C	SER	A	184	32.557	2.896	12.675	1.00	32.25	C	
ANISOU	1315	C	SER	A	184	3525	4901	3827	255	-164	454	C
ATOM	1316	O	SER	A	184	31.344	2.392	12.468	1.00	32.73	O	
ANISOU	1316	O	SER	A	184	3524	5047	3874	263	-194	472	O
ATOM	1317	CB	SER	A	184	32.620	0.281	13.181	1.30	32.50	C	
ANISOU	1317	CB	SER	A	184	3506	4347	3805	139	-230	402	C
ATOM	1318	OG	SER	A	184	32.854	0.655	14.564	1.00	31.97	O	
ANISOU	1318	OG	SER	A	184	3491	4809	3847	86	-185	414	O
ATOM	1319	N	GLY	A	185	33.320	9.541	19.384	1.00	34.61	N	
ANISOU	1319	N	GLY	A	185	3349	5140	4160	2781	-105	480	N
ATOM	1320	CA	GLY	A	185	32.736	4.657	14.143	1.00	29.59	C	
ANISOU	1320	CA	GLY	A	185	3170	4523	3548	320	-71	505	C
ATOM	1321	C	GLY	A	185	31.551	4.246	14.987	1.00	32.76	C	
ANISOU	1321	C	GLY	A	185	3493	4990	3959	275	-93	495	C
ATOM	1322	O	GLY	A	185	30.661	5.075	15.211	1.09	33.43	O	
ANISOU	1322	O	GLY	A	185	3535	5133	4053	322	-83	512	O
ATOM	1323	N	SER	A	186	31.524	2.993	15.479	1.00	36.20	N	
ANISOU	1323	N	SER	A	186	3929	5424	4401	168	-120	458	N
ATOM	1324	CA	SER	A	186	30.354	2.494	16.225	1.00	32.53	C	
ANISOU	1324	CA	SER	A	186	3392	5027	3942	184	-140	459	C
ATOM	1325	C	SER	A	186	30.751	1.851	17.540	1.00	34.48	C	
ANISOU	1325	C	SER	A	186	3652	5231	4219	59	-121	448	C
ATOM	1326	O	SER	A	186	29.935	1.207	18.162	1.06	34.41	O	

TABLE 3-continued

ANISOU	1326	O	SER	A	186	3593	5259	4211	-5	-133	432	O
ATOM	1327	CB	SER	A	186	29.494	1.530	15.372	1.00	32.76	C	
ANISOU	1327	CB	SER	A	186	3387	5121	3938	92	-207	445	C
ATOM	1328	OG	SER	A	186	29.137	2.086	14.036	1.00	32.91	O	
ANISOU	1328	OG	SER	A	186	3401	5184	3920	135	-293	456	O
ATOM	1329	N	THR	A	187	32.305	2.023	17.958	1.00	29.66	N	
ANISOU	1329	N	THR	A	187	3104	4534	13630	55	-88	446	N
ATOM	1330	CA	THR	A	187	32.472	1.423	19.230	1.00	32.19	C	
ANISOU	1330	CA	THA	A	187	3444	4810	5976	0	-74	435	C
ATOM	1331	C	THR	A	187	32.107	2.299	20.421	1.00	32.59	C	
ANISOU	1331	C	THR	A	187	3459	4331	4044	18	-27	447	C
ATOM	1332	O	THR	A	187	32.475	3.440	20.509	1.00	33.52	O	
ANISOU	1332	O	THR	A	187	3208	4598	3795	85	13	456	O
ATOM	1333	CB	THR	A	187	33.994	1.188	19.191	1.00	35.12	C	
ANISOU	1333	CB	THR	A	187	3894	5034	41365	-1	-65	425	C
ATOM	1334	OG1	THR	A	187	34.255	0.259	18.115	1.00	41.03	O	
ANISOU	1334	OG1	THR	A	187	4573	5320	5095	-20	-110	406	O
ATOM	1335	OG2	THR	A	187	34.443	0.515	23.498	1.00	34.33	C	
ANISOU	1335	OG2	THR	A	187	3515	4940	4289	-67	-50	417	C
ATOM	1336	N	LYS	A	188	31.364	1.749	21.363	1.00	33.54	N	
ANISOU	1336	N	LYS	A	188	3537	5043	4162	-45	-30	445	N
ATOM	1337	CA	LYS	A	188	30.557	2.571	22.304	1.00	33.53	C	
ANISOU	1337	CA	LYS	A	188	3492	5100	4175	-24	12	453	C
ATOM	1338	C	LYS	A	188	31.673	3.519	23.152	1.00	35.82	C	
ANISOU	1338	C	LYS	A	188	3814	5313	4479	20	64	451	C
ATOM	1339	O	LYS	A	188	31.227	4.744	23.246	1.00	30.97	O	
ANISOU	1339	O	LYS	A	188	3174	4721	3871	94	99	455	O
ATOM	1340	CB	LYS	A	188	29.854	1.528	23.180	1.00	34.71	C	
ANISOU	1340	CB	LYS	A	188	3590	5292	4396	-115	2	451	C
ATOM	1341	CG	LYS	A	188	28.645	2.207	23.358	1.00	48.11	C	
ANISOU	1341	CG	LYS	A	188	5201	7082	5995	-104	33	455	C
ATOM	1342	CD	LYS	A	188	29.051	2.493	25.263	1.00	45.16	C	
ANISOU	1342	CD	LYS	A	188	5229	7062	5039	-118	80	453	C
ATOM	1343	CE	LYS	A	188	27.950	2.161	25.263	1.00	48.58	C	
ANISOU	1343	CE	LYS	A	188	5211	7201	6346	-177	100	456	C
ATOM	1344	NZ	LYS	A	188	28.389	2.901	27.512	1.00	47.47	N	
ANISOU	1344	NZ	LYS	A	188	5092	7037	5907	-152	155	450	N
ATOM	1345	N	GLU	A	189	82.502	2.961	23.798	1.00	28.14	N	
ANISOU	1345	N	GLU	A	189	2905	4278	3524	-25	55	443	N
ATOM	1346	CA	GLU	A	189	83.419	3.772	24.685	1.00	84.41	C	

TABLE 3-continued

ANISOU	1346	CA	GLU	A	189	3726	5014	4334	5	107	433	C
ATOM	1347	C	GLU	A	189	34.341	4.713	23.872	1.00	30.66	C	
ANISOU	1347	C	GLU	A	189	3319	4503	3905	82	123	439	C
ATOM	1348	O	GLU	A	189	34.595	5.835	24.299	1.00	29.73	O	
ANISOU	1348	O	GLU	A	189	3135	4331	3781	135	163	437	O
ATOM	1349	CB	GLU	A	189	34.165	2.921	25.770	1.00	29.82	C	
ANISOU	1349	CB	GLU	A	189	3191	4261	3757	-65	101	430	C
ATOM	1350	CG	GLU	A	189	33.132	2.312	26.709	1.00	31.20	C	
ANISOU	1350	CG	GLU	A	189	3222	4524	3909	-132	101	435	C
ATOM	1351	CD	GLU	A	189	33.679	1.700	27.995	1.00	34.59	C	
ANISOU	1351	CD	GLU	A	189	3792	5015	4334	-194	103	434	C
ATOM	1352	OE1	GLU	A	189	32.921	1.139	28.803	1.00	33.24	O	
ANISOU	1352	OE1	GLU	A	189	3594	4896	4140	-257	106	444	O
ATOM	1353	OE2	GLU	A	189	34.884	1.761	23.245	1.00	38.15	O	
ANISOU	1353	OE2	GLU	A	189	4303	5383	4804	-184	102	425	O
ATOM	1354	N	VAL	A	190	34.644	4.327	22.543	1.00	29.85	N	
ANISOU	1354	N	VAL	A	190	3211	4359	3770	90	04	443	N
ATOM	1355	CA	VAL	A	190	35.447	5.246	21.813	1.00	28.36	C	
ANISOU	1355	CA	VAL	A	190	3135	4198	3672	159	115	450	C
ATOM	1356	C	VAL	A	190	34.501	5.457	21.472	1.00	33.83	C	
ANISOU	1356	C	VAL	A	190	3711	4359	4282	234	137	457	C
ATOM	1357	O	VAL	A	190	35.103	7.508	21.605	1.00	33.48	O	
ANISOU	1357	O	VAL	A	190	3639	4771	4262	289	174	479	O
ATOM	1358	CB	VAL	A	190	35.938	4.550	20.534	1.30	28.08	C	
ANISOU	1358	CB	VAL	A	190	3055	4053	3546	153	82	449	C
ATOM	1359	CG1	VAL	A	190	35.568	5.533	49.564	1.39	27.77	C	
ANISOU	1359	CG1	VAL	A	190	3045	3994	3511	224	137	464	C
ATOM	1360	CG2	VAL	A	190	37.019	3.540	20.978	1.50	27.25	C	
ANISOU	1360	CG2	VAL	A	190	2987	3898	3457	95	56	428	C
ATOM	1361	N	ILE	A	191	33.382	6.261	20.944	1.00	32.29	N	
ANISOU	1361	N	ILE	A	191	3462	4745	4551	239	111	475	N
ATOM	1362	CA	ILE	A	191	32.412	7.370	20.618	1.00	31.82	C	
ANISOU	1362	CA	ILE	A	191	3354	4740	3995	316	124	492	C
ATOM	1363	C	ILE	A	191	32.173	8.268	21.788	1.00	34.76	C	
ANISOU	1363	C	ILE	A	191	3705	5112	4339	345	167	485	C
ATOM	1364	O	ILE	A	191	32.057	9.477	21.531	1.00	36.34	O	
ANISOU	1364	O	ILE	A	191	3905	5301	4603	423	193	495	O
ATOM	1365	CB	ILE	A	191	30.999	3.781	20.340	1.00	34.40	C	
ANISOU	1365	CB	ILE	A	191	3606	5170	4293	295	86	493	C
ATOM	1366	CG1	ILE	A	191	30.971	6.064	19.009	1.00	40.95	C	

TABLE 3-continued

ANISOU	1366	CG1	ILE	A	191	4448	6016	5095	283	37	438	C
ATOM	1367	CG2	ILE	A	191	29.879	7.841	20.373	1.00	44.85	C	
ANISOU	1367	CG2	ILE	A	191	4866	6561	5618	369	100	502	C
ATOM	1368	CD1	ILE	A	191	23.541	5.453	18.906	1.00	40.26	C	
ANISOU	1368	CD1	ILE	A	191	4279	6033	4986	251	-2	493	C
ATOM	1369	N	HIS	A	192	32.055	7.551	22.088	1.00	30.94	N	
ANISOU	1369	N	HIS	A	192	3206	4643	3907	281	175	457	N
ATOM	1370	CA	HIS	A	192	31.707	8.417	24.165	1.00	31.50	C	
ANISOU	1370	CA	HIS	A	192	3239	4766	4027	303	216	454	C
ATOM	1371	C	HIS	A	192	32.835	9.394	24.511	1.00	31.70	C	
ANISOU	1371	C	HIS	A	192	3337	4560	4046	346	252	448	C
ATOM	1372	O	HIS	A	192	32.508	10.595	24.771	1.00	27.87	O	
ANISOU	1372	O	HIS	A	192	2843	4170	3578	416	284	445	O
ATOM	1373	CB	HIS	A	192	31.399	7.465	25.317	1.00	30.59	C	
ANISOU	1373	CB	HIS	A	192	3114	4649	3860	218	215	440	C
ATOM	1174	CG	HIS	A	192	31.146	8.190	26.507	1.00	30.95	C	
ANISOU	1374	CG	HIS	A	192	3143	4703	3910	237	262	422	C
ATOM	1375	ND1	HIS	A	192	30.014	8.974	25.816	1.00	29.94	N	
ANISOU	1375	ND1	HIS	A	192	2945	4655	3777	232	285	417	N
ATOM	1376	CD2	HIS	A	192	31.914	3.273	27.85	1.00	29.67	C	
ANISOU	1376	CD2	HIS	A	192	3021	4497	3755	212	263	405	C
ATOM	1377	CE1	HIS	A	192	30.085	9.526	28.028	1.00	23.35	C	
ANISOU	1377	CE1	HIS	A	192	2745	4448	3078	303	327	395	C
ATOM	1378	NE2	HIS	A	192	31.254	9.933	28.621	1	00	33.25	N
ANISOU	1378	NE2	HIS	A	192	3056	4614	3825	253	327	388	N
ATOM	1379	N	ALA	A	193	34.571	8.909	24.482	1.00	29.31	N	
ANISOU	1379	N	ALA	A	193	3096	4234	3757	307	245	444	N
ATOM	1380	CA	ALA	A	193	36.181	9.700	24.752	1.00	33.59	C	
ANISOU	1380	CA	ALA	A	193	3312	4350	3052	338	274	437	C
ATOM	1381	C	ALA	A	193	35.299	10	815	23.626	1.00	26.36	C
ANISOU	1381	C	ALA	A	193	2353	3358	3491	417	285	480	C
ATOM	1382	O	ALA	A	193	35.453	12.041	23.934	1.00	30.25	O	
ANISOU	1382	O	ALA	A	193	3297	4251	3549	474	319	458	O
ATOM	1383	CB	ALA	A	193	35.496	8.923	24.940	3	00	24.05	C
ANISOU	1383	CB	ALA	A	193	2538	3454	3137	277	250	425	C
ATOM	1384	N	ALA	A	194	35.156	10.433	22.351	1.00	30.88	N	
ANISOU	1384	N	ALA	A	194	2364	4388	3982	424	258	482	N
ATOM	1385	CA	ALA	A	194	35.313	13.460	21.304	1.00	39.34	C	
ANISOU	1385	CA	ALA	A	194	3310	4289	3919	490	270	510	C
ATOM	1386	C	ALA	A	194	34.243	12.539	21.476	1.00	34.26	C	

TABLE 3-continued

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ANISOU	1386	C	ALA	A	194	3775	4825	4413	571	234	518	C
ATOM	1387	O	ALA	A	194	34.490	13.754	21.329	1.00	34.92	O	
ANISOU	1387	O	ALA	A	194	3885	4857	4521	637	332	532	O
ATOM	1388	CB	ALA	A	194	35.236	10.865	19.872	1.09	31.58	C	
ANISOU	1388	CB	ALA	A	194	3482	4473	4043	448	235	533	C
ATOM	1389	N	GLY	A	195	33.033	12.031	21.792	1.00	31.93	N	
ANISOU	1389	N	GLY	A	195	3413	4620	4100	561	268	510	N
ATOM	1390	CA	GLY	A	195	31.833	12.939	22.059	1.00	32.14	C	
ANISOU	1390	CA	GLY	A	195	3381	4704	4125	628	275	509	C
ATOM	1391	C	GLY	A	195	32.027	13.933	23.162	1.00	35.94	C	
ANISOU	1391	C	GLY	A	195	3520	4388	4337	653	320	467	C
ATOM	1392	O	GLY	A	195	31.552	15.057	23.377	1.00	36.40	O	
ANISOU	1392	O	GLY	A	195	3922	5195	4714	748	335	493	O
ATOM	1393	N	LEU	A	196	32.774	13.559	24.199	1.00	33.73	N	
ANISOU	1393	N	LEU	A	196	3620	4823	4372	605	338	461	N
ATOM	1394	CA	LEU	A	196	33.072	14.562	25.256	1.00	32.59	C	
ANISOU	1394	CA	LEU	A	196	3532	4655	4295	637	373	434	C
ATOM	1395	C	LEU	A	196	33.913	15.732	24.321	1.00	31.63	C	
ANISOU	1395	C	LEU	A	199	3450	4404	4201	697	399	446	C
ATOM	1396	O	LEU	A	196	33.599	16.941	25.256	1.00	32.02	O	
ANISOU	1396	O	LEU	A	196	3482	4436	4247	703	425	434	O
ATOM	1397	CB	LEU	A	196	33.791	13.853	25.328	1.00	30.61	C	
ANISOU	1397	CB	LEU	A	196	3250	4353	4005	559	383	404	C
ATOM	1398	CG	LEU	A	196	32.776	12.922	27.172	1.00	34.22	C	
ANISOU	1398	CG	LEU	A	196	3059	4316	4426	507	379	350	C
ATOM	1399	CD1	LEU	A	196	33.621	11.839	27.992	1.00	30.65	C	
ANISOU	1399	CD1	LEU	A	196	3242	4437	3966	414	371	376	C
ATOM	1400	CD2	LEU	A	196	31.776	13.740	28.034	1.00	36.84	C	
ANISOU	1400	CD2	LEU	A	196	3940	5302	4755	559	412	365	C
ATOM	1401	N	ALA	A	197	34.935	15.551	23.994	1.00	31.49	N	
ANISOU	1401	N	ALA	A	197	3468	4330	4155	573	390	469	N
ATOM	1402	CA	ALA	A	157	35.745	16.613	23.425	1.00	32.73	C	
ANISOU	1402	CA	ALA	A	197	3682	4396	4357	719	410	490	C
ATOM	1403	C	ALA	A	197	34.395	17.415	22.429	1.50	38.15	C	
ANISOU	1403	C	ALA	A	197	4355	5103	5035	334	434	527	C
ATOM	1404	O	ALA	A	197	34.857	13.650	22.471	1.00	33.58	O	
ANISOU	1404	O	ALA	A	197	3338	4510	4525	372	425	533	O
ATOM	1405	CB	ALA	A	197	35.954	16.031	22.792	1.00	31.28	C	
ANISOU	1405	CB	ALA	A	197	3546	4152	4177	659	404	536	C
ATOM	1406	N	TYR	A	198	34.218	16.703	21	534	1.00	36.44	N

TABLE 3-continued

ANISOU	1406	N	TYR	A	198	4106	4955	4778	802	360	551	N
ATOM	1407	CA	TYR	A	198	33.351	17.353	20.595	1.00	43.04	C	
ANISOU	1407	CA	TYR	A	198	4572	5473	5246	882	354	586	C
ATOM	1408	C	TYR	A	198	32.332	18.207	21.276	1.00	40.17	C	
ANISOU	1408	C	TYR	A	198	4522	5486	5254	955	367	559	C
ATOM	1409	O	TYR	A	198	32.193	19.440	20.822	1.00	44.53	O	
ANISOU	1409	O	TYR	A	198	5095	5997	5325	1039	374	594	O
ATOM	1410	CB	TYR	A	198	32.557	16.334	19.785	1.00	43.37	C	
ANISOU	1410	CB	TYR	A	198	4922	5962	5593	861	309	600	C
ATOM	1411	CG	TYR	A	198	31.045	16.964	18.515	1.00	50.57	C	
ANISOU	1411	CG	TYR	A	198	5527	6904	6465	945	285	642	C
ATOM	1412	CD1	TYR	A	198	32.518	17.199	17.339	1.00	52.24	C	
ANISOU	1412	CD1	TYR	A	198	6090	7067	5531	952	280	687	C
ATOM	1413	CD2	TYR	A	198	30.482	17.203	18.718	1.00	50.02	C	
ANISOU	1413	CD2	TYR	A	198	5686	5913	6436	1036	256	539	C
ATOM	1414	CE1	TYR	A	198	31.858	17.760	16.303	1.00	53.65	C	
ANISOU	1414	CE1	TYR	A	198	6282	7275	5838	1038	254	733	C
ATOM	1415	CE2	TYR	A	198	29.780	17.357	17.635	1.00	55.84	C	
ANISOU	1415	CE2	TYR	A	198	5782	7044	6430	1337	236	670	C
ATOM	1416	CZ	TYR	A	198	30.477	18.071	10.427	1.00	56.52	C	
ANISOU	1416	CZ	TYR	A	198	6795	7941	7413	1102	220	726	C
ATOM	1417	OH	TYR	A	198	29.850	18.594	15.320	1.00	63.59	O	
ANISOU	1417	OH	TYR	A	198	7468	8535	8057	1181	194	770	O
ATOM	1418	N	LYS	A	199	31.318	17.374	22.435	1.00	42.42	N	
ANISOU	1418	N	LYS	A	199	4754	5829	5536	927	374	527	N
ATOM	1419	CA	LYS	A	199	30.840	18.723	23.113	1.00	46.20	C	
ANISOU	1419	CA	LYS	A	199	4813	5057	5555	939	390	503	C
ATOM	1420	C	LYS	A	199	21.426	19.943	20.823	1.00	46.02	C	
ANISOU	1420	C	LYS	A	199	5215	6220	6352	1044	429	462	C
ATOM	1421	O	LYS	A	199	30.737	20.029	24.112	1.00	39.55	O	
ANISOU	1421	O	LYS	A	199	4389	5413	5263	1128	442	469	O
ATOM	1422	CB	LYS	A	199	20.914	17.912	24.030	1.00	46.72	C	
ANISOU	1422	CB	LYS	A	199	5171	6513	5009	958	389	466	C
ATOM	1423	CG	LYS	A	199	29.302	17.001	23.224	1.00	48.71	C	
ANISOU	1423	CG	LYS	A	199	5350	6357	6251	940	1347	457	C
ATOM	1424	CD	LYS	A	199	28.429	15.852	24.056	1.00	59.06	C	
ANISOU	1424	CD	LYS	A	199	6720	8386	7682	358	345	458	C
ATOM	1425	CE	LYS	A	199	27.888	14.737	23.157	1.00	61.96	C	
ANISOU	1425	CE	LYS	A	199	1931	8715	7901	811	297	479	C
ATOM	1426	NZ	LYS	A	199	26.594	14.101	23.536	1.00	57.90	N	

TABLE 3-continued

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ANISOU 1426 NZ LYS A 199	6314 3323 7304 787 259 460 N
ATOM 1427 N ARG A 200	32.721 19.032 24.020 1.00 40.56 N
ANISOU 1427 N ARG A 200	4592 5441 5384 593 445 473 N
ATOM 1428 CA ARG A 200	33.361 21.087 24.606 1.00 39.35 C
ANISOU 1428 CA ARG A 200	4491 5135 5275 1028 478 469 C
ATOM 1429 C ARG A 200	34.005 21.968 23.532 1.00 39.25 C
ANISOU 1429 C ARG A 200	4545 5078 5291 1070 480 507 C
ATOM 1430 O ARG A 200	34.880 22.752 23.689 1.00 37.91 O
ANISOU 1430 O ARG A 200	4433 4807 5164 1072 505 497 O
ATOM 1431 CB ARG A 200	34.455 20.509 25.531 1.00 37.57 C
ANISOU 1431 CB ARG A 200	4295 4015 5062 544 494 424 111 C
ATOM 1432 CG ARG A 200	33.390 20.115 26.825 1.00 33.23 C
ANISOU 1432 CG ARG A 200	4464 5232 5257 915 592 373 C
ATOM 1433 CD ARG A 200	34.974 19.404 27 556 1.00 43.31 C
ANISOU 1433 CD ARG A 200	5007 5530 5770 625 505 349 C
ATOM 1434 NE ARG A 200	36.028 20.334 27.995 1.00 45.27 N
ANISOU 1434 NE ARG A 200	5319 5817 6065 834 527 328 N
ATOM 1435 CZ ARG A 200	35.914 21.139 29.050 1.00 41.30 C
ANISOU 1435 CZ ARG A 200	4823 5290 5560 366 550 281 C
ATOM 1436 NH1 ARG A 200	34.798 21.161 29.783 1.00 46.48 N
ANISOU 1436 NH1 ARG A 200	5426 6327 6209 808 560 250 N
ATOM 1437 NH2 ARG A 200	36.932 21.900 29.384 1.00 41.85 N
ANISOU 1437 NH2 ARG A 200	4950 5255 5694 364 554 261 N
ATOM 1438 N ASP A 201	33.581 21.826 22.286 1.00 36.25 N
ANISOU 1438 N ASP A 201	4153 4731 4884 1100 454 558 N
ATOM 1439 CA ASP A 201	34.106 22.599 21.142 1.00 37.99 C
ANISOU 1439 CA ASP A 201	4443 4872 5119 1140 455 615 C
ATOM 1440 C ASP A 201	35.616 22.445 20.890 1.00 41.36 C
ANISOU 1440 C ASP A 201	4999 5276 5628 1069 476 628 C
ATOM 1441 O ASP A 201	36.254 23.377 20.410 1.00 35.40 O
ANISOU 1441 O ASP A 201	4244 4365 4841 1095 495 660 O
ATOM 1442 CB ASP A 201	33.719 24.079 21.264 1.00 43.95 C
ANISOU 1442 CB ASP A 201	5223 5562 5913 1239 469 621 C
ATOM 1443 CG ASP A 201	32.249 24.217 21.525 1.00 46 19 C
ANISOU 1443 CG ASP A 201	5434 5936 6181 1313 450 503 C
ATOM 1444 OD1 ASP A 201	31.479 23.656 20.725 1.00 48.93 O
ANISOU 1444 OD1 ASP A 201	5737 6371 6485 1328 415 631 O
ATOM 1445 OD2 ASP A 201	31.849 24.772 22.567 1.00 50.88 O
ANISOU 1445 OD2 ASP A 201	6007 6525 6800 1349 469 554 O
ATOM 1446 N ILE A 202	36.157 21.255 21.194 1.00 34.53 N

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TABLE 3-continued

ANISOU	1446	N	ILE	A	202	4055	4082	4681	981	470	604	N
ATOM	1447	CA	ILE	A	202	37.569	20.956	20.807	1.00	35.66	C	
ANISOU	1447	CA	ILE	A	202	4250	4459	4841	914	486	513	C
ATOM	1448	C	ILE	A	202	37.532	20.153	19.628	1.00	33.46	C	
ANISOU	1448	C	ILE	A	202	3969	4230	4515	899	463	654	C
ATOM	1449	O	ILE	A	202	36.662	19.310	19.530	1.00	37.38	O	
ANISOU	1449	O	ILE	A	202	4413	4819	4969	892	431	647	O
ATOM	1450	CB	ILE	A	202	38.170	20.126	22.056	1.00	31.89	C	
ANISOU	1450	CB	ILE	A	202	3756	3987	4372	6134	490	558	C
ATOM	1451	CG1	ILE	A	202	38.051	20.941	23.367	1.00	34.15	C	
ANISOU	1451	CG1	ILE	A	202	4043	4236	4596	857	510	511	C
ATOM	1452	CG2	ILE	A	202	39.623	19.658	21.718	1.00	28.60	C	
ANISOU	1452	CG2	ILE	A	202	3379	3514	3970	765	500	563	C
ATOM	1453	CD1	ILE	A	202	38.546	22.395	20.321	1.00	36.33	C	
ANISOU	1453	CD1	ILE	A	202	4375	4404	5026	905	538	521	C
ATOM	1454	N	PRO	A	203	38.376	20.462	18.607	1.00	36.85	N	
ANISOU	1454	N	PRO	A	203	4462	4603	4946	895	478	697	N
ATOM	1455	CA	PRO	A	203	38.280	19.689	17.383	1.00	32.80	C	
ANISOU	1455	CA	PRO	A	203	3938	4146	4379	885	455	732	C
ATOM	1456	C	PRO	A	203	36.545	18.172	17.522	1.00	34.34	C	
ANISOU	1456	C	PRO	A	203	4132	4399	4546	307	433	698	C
ATOM	1457	O	PRO	A	203	39.423	17	718	18.312	1.00	29.30	O
ANISOU	1457	O	PRO	A	203	3468	3728	3936	745	447	560	O
ATOM	1458	CB	PRO	A	203	39.364	20.312	16.437	1.00	36.68	C	
ANISOU	1458	CB	PRO	A	203	4498	4557	4881	884	487	779	C
ATOM	1459	CG	PRO	A	203	40.360	20.893	17.426	1.00	41.83	C	
ANISOU	1459	CG	PRO	A	203	5175	5117	5600	851	525	749	C
ATOM	1460	CD	PRO	A	203	39.492	21.430	18.561	1.00	41.64	C	
ANISOU	1460	CD	PRO	A	203	5121	5098	5602	891	518	713	C
ATOM	1461	N	VAL	A	204	37.775	17.401	16.759	1.00	28.37	N	
ANISOU	1461	N	VAL	A	204	3316	3727	3735	813	395	712	N
ATOM	1462	CA	VAL	A	204	37.597	15.935	16.776	1.00	28.92	C	
ANISOU	1462	CA	VAL	A	204	3361	3852	3776	744	368	682	C
ATOM	1463	C	VAL	A	204	38.044	15.529	5.359	1.00	29.16	C	
ANISOU	1463	C	VAL	A	204	3414	3907	3758	748	353	715	C
ATOM	1464	O	VAL	A	204	37.164	15.886	14.568	1.00	29.73	O	
ANISOU	1464	O	VAL	A	204	3478	4023	3701	805	321	749	O
ATOM	1465	CB	VAL	A	204	36.302	15.267	17.354	1.00	30.57	C	
ANISOU	1465	CB	VAL	A	204	3501	4150	3964	737	331	354	C
ATOM	1466	CG1	VAL	A	204	36.580	13.712	17.293	1.00	29.25	C	

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ANISOU	1466	CG1	VAL	A	204	3313	4035	3765	864	297	629	C
ATOM	1467	CG2	VAL	A	204	33.425	15.701	18.850	1.00	23.94	C	
ANISOU	1467	CG2	VAL	A	204	3271	3927	3798	737	351	613	C
ATOM	1468	N	VAL	A	205	39.055	14.745	15.035	1.00	28.03	N	
ANISOU	1468	N	VAL	A	205	3295	3746	3609	693	359	703	N
ATOM	1469	CA	VAL	A	205	39.282	14.245	13.635	1.00	27.42	C	
ANISOU	1469	CA	VAL	A	205	3242	8698	3477	695	345	728	C
ATOM	1470	C	VAL	A	205	39.057	12.709	13.704	1.00	31.77	C	
ANISOU	1470	C	VAL	A	205	3762	4309	4001	637	303	688	C
ATOM	1471	O	VAL	A	205	39.705	12.045	14.504	1.00	33.42	O	
ANISOU	1471	O	VAL	A	205	3933	4492	4239	581	333	647	O
ATOM	1472	CB	VAL	A	205	40.714	14.506	13.145	1.00	27.28	C	
ANISOU	1472	CB	VAL	A	205	3273	3605	3470	675	353	741	C
ATOM	1473	CG1	VAL	A	205	41.000	13.875	11.755	1.00	28.33	C	
ANISOU	1473	CG1	VAL	A	205	3437	3780	3547	672	364	758	C
ATOM	1474	CG2	VAL	A	205	40.949	16.046	13.028	1.00	31.16	C	
ANISOU	1474	CG2	VAL	A	205	3308	4033	3998	725	436	788	C
ATOM	1475	N	SER	A	206	38.190	12.157	12.873	1.00	32.33	N	
ANISOU	1475	N	SER	A	206	3814	4455	4015	551	259	394	N
ATOM	1476	CA	SER	A	206	37.973	10.713	12.900	1.00	34.46	C	
ANISOU	1476	CA	SER	A	206	4059	4774	4261	594	213	654	C
ATOM	1477	C	SER	A	206	38.725	10.053	11.775	1.00	35.82	C	
ANISOU	1477	C	SER	A	206	4271	4945	4394	579	213	552	C
ATOM	1478	O	SER	A	206	38.715	10.571	10.647	1.00	25.54	O	
ANISOU	1478	O	SER	A	206	3517	4175	3570	5624	219	589	O
ATOM	1479	CB	SER	A	206	36.493	10.413	12.710	1.00	34.89	C	
ANISOU	1479	CB	SER	A	206	4060	4916	4279	611	164	855	C
ATOM	1480	OG	SER	A	206	35.931	10.754	13.542	1.00	42.02	O	
ANISOU	1480	OG	SER	A	206	4921	5822	5221	309	172	843	O
ATOM	1481	N	LEU	A	207	39.333	8.520	12.096	1.00	31.44	N	
ANISOU	1481	N	LEU	A	207	3717	4378	3849	518	201	308	N
ATOM	1482	CA	LEU	A	207	39.970	8.105	11.003	1.00	32.95	C	
ANISOU	1482	CA	LEU	A	207	3945	4583	4033	502	192	594	C
ATOM	1483	C	LEU	A	207	39.255	5.789	10.972	1.00	33.08	C	
ANISOU	1483	C	LEU	A	207	3931	4550	3937	462	130	553	C
ATOM	1484	O	LEU	A	207	39.400	5.985	11.882	1.00	31.37	O	
ANISOU	1484	O	LEU	A	207	3700	4417	3804	409	114	522	O
ATOM	1485	CB	LEU	A	207	41.395	7.824	11.423	1.00	33.46	C	
ANISOU	1485	CB	LEU	A	207	4032	4576	4106	467	228	568	C
ATOM	1486	CG	LEU	A	207	42.367	8.951	11.583	1.00	36.75	C	

TABLE 3-continued

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ANISOU	1486	CG	LEU	A	207	4475	4525	4563	489	293	595	C
ATOM	1487	CD1	LEU	A	207	43.802	8.449	11.673	1.00	37.87	0	C
ANISOU	1487	CD1	LEU	A	207	4538	5018	4734	452	320	562	C
ATOM	1488	CD2	LEU	A	207	42.284	9.979	10.389	1.00	36.79	0	C
ANISOU	1488	CD2	LEU	A	207	4511	4941	4525	545	320	850	C
ATOM	1489	N	THR	A	208	38.455	6.544	9.900	1.00	33.55	0	N
ANISOU	1489	N	THR	A	208	4000	4783	3997	485	90	567	N
ATOM	1490	CA	THR	A	208	37.541	5.357	9.909	1.00	32.07	0	C
ANISOU	1490	CA	THR	A	208	3755	4647	3774	443	24	532	C
ATOM	1491	C	THR	A	208	37.534	4.725	8.545	1.00	32.35	0	C
ANISOU	1491	C	THR	A	208	3831	4716	3744	451	-6	519	C
ATOM	1492	O	THR	A	208	38.458	5.135	7.712	1.00	31.69	0	O
ANISOU	1492	O	THR	A	208	3794	452	3635	484	29	535	O
ATOM	1493	CB	THR	A	208	36.049	5.831	9.980	1.00	39.50	0	C
ANISOU	1493	CB	THR	A	208	4652	5653	4700	472	-10	556	C
ATOM	1494	OG1	THR	A	208	35.754	6.512	8.729	1.00	33.53	0	O
ANISOU	1494	OG1	THR	A	208	4543	5571	4520	537	-15	592	O
ATOM	1495	CG2	THR	A	208	35.899	6.330	11.034	1.00	36.41	0	C
ANISOU	1495	CG2	THR	A	208	4490	5403	4011	491	27	577	C
ATOM	1496	N	ASN	A	209	36.808	3.712	8.324	1.00	29.02	0	N
ANISOU	1496	N	ASN	A	209	3423	4386	3332	417	-70	486	N
ATOM	1497	CA	ASN	A	209	36.599	3.100	7.005	1.00	23.54	0	C
ANISOU	1497	CA	ASN	A	209	3350	4333	3154	425	-109	471	C
ATOM	1498	C	ASN	A	209	35.245	3.122	6.568	1.00	32.35	0	C
ANISOU	1498	C	ASN	A	209	3850	4902	3669	441	-169	481	C
ATOM	1499	O	ASN	A	209	34.533	2.307	5.753	1.00	33.31	0	O
ANISOU	1499	O	ASN	A	209	3912	5067	3678	425	-224	452	O
ATOM	1500	CB	ASN	O	209	37.149	1.582	7.087	1.00	25.05	0	C
ANISOU	1500	CB	ASN	A	209	3413	4359	2223	359	-143	407	C
ATOM	1501	CG	ASN	A	209	38.628	1.420	7.407	1.00	32.49	0	C
ANISOU	1501	CG	ASN	A	209	3909	4723	3709	343	-92	389	C
ATOM	1502	OD1	ASN	A	209	39.411	0.963	6.561	1.00	32.52	0	O
ANISOU	1502	OD1	ASN	A	209	3968	4733	3693	355	-86	363	O
ATOM	1503	ND2	ASN	A	209	35.997	1.374	3.627	1.00	25.74	0	N
ANISOU	1503	ND2	ASN	A	209	3039	3822	2920	226	-52	395	N
ATOM	1504	N	ILE	A	210	34.483	4.107	7.046	1.00	31.45	0	N
ANISOU	1504	N	ILE	A	210	3631	4408	3511	475	-150	521	N
ATOM	1505	CA	ILE	A	210	32.126	4.238	6.586	1.00	37.91	0	C
ANISOU	1505	CA	ILE	A	210	4430	5712	4292	501	-214	533	C
ATOM	1506	C	ILE	A	210	32.725	5.739	60.601	1	00	34.78	C

TABLE 3-continued

ANISOU	1506	C	ILE	A	210	3990	5325	3930	581	-135	592	C
ATOM	1507	O	ILE	A	210	33.123	5.475	7.480	1.00	37.23	O	
ANISOU	1507	O	ILE	A	210	4297	5578	4260	592	-132	511	O
ATOM	1508	CB	ILE	A	210	32.179	3.303	7.402	1.00	38.91	C	
ANISOU	1508	CB	ILE	A	210	4460	5877	4448	432	-293	497	C
ATOM	1509	CG1	ILE	A	210	30.869	3.133	6.575	1.00	42.64	C	
ANISOU	1509	CG1	ILE	A	210	4882	6446	4873	445	-333	495	C
ATOM	1510	CG2	ILE	A	210	31.992	3.775	8.733	1.00	36.54	C	
ANISOU	1510	CG2	ILE	A	210	4117	5353	4212	419	-227	509	C
ATOM	1511	CD1	ILE	A	210	30.729	1.731	6.144	1.00	42.06	C	
ANISOU	1511	CD1	ILE	A	210	481.5	6395	4760	381	-395	442	C
ATOM	1512	N	ASN	A	211	32.335	6.171	5.567	1.00	37.89	N	
ANISOU	1512	N	ASN	A	211	4381	5780	4235	639	-221	619	N
ATOM	1513	CA	ASN	A	211	31.559	7.555	5.504	1.00	41.85	C	
ANISOU	1513	CA	ASN	A	211	4373	6291	4734	722	-203	676	C
ATOM	1514	C	ASN	A	211	30.559	7.999	6.535	1.00	40.65	C	
ANISOU	1514	C	ASN	A	211	4645	6169	4640	732	-239	650	C
ATOM	1515	O	ASN	A	211	30.654	9.131	7.002	1.00	43.30	O	
ANISOU	1515	O	ASN	A	211	5046	6527	5070	783	-166	716	O
ATOM	1516	CB	ASN	A	211	31.073	7.872	4.079	1.00	41.05	C	
ANISOU	1516	CB	ASN	A	211	4792	6251	4553	785	-247	706	C
ATOM	1517	CG	ASN	A	211	32.217	7.817	3.114	1.00	45.85	C	
ANISOU	1517	CG	ASN	A	211	5484	6821	5111	792	-216	717	C
ATOM	1518	OD1	ASN	A	211	33.112	5.648	3.173	1.00	48.31	O	
ANISOU	1518	OD1	ASN	A	211	5849	7065	5441	819	-149	754	O
ATOM	1519	ND2	ASN	A	211	32.293	6.751	2.349	1.00	52.29	N	
ANISOU	1519	ND2	ASN	A	211	6321	7673	5573	757	-257	670	N
ATOM	1520	N	HIS	A	212	29.590	7.139	6.879	1.00	35.59	N	
ANISOU	1520	N	HIS	A	212	4313	5972	4380	684	-252	643	N
ATOM	1521	CA	HIS	A	212	2.575	7.558	7.450	1.00	35.54	C	
ANISOU	1521	CA	HIS	A	212	3853	5637	4051	694	-285	645	C
ATOM	1522	C	HIS	A	212	28.474	6.634	9.018	1.00	34.41	C	
ANISOU	1522	C	HIS	A	212	3854	5479	3942	504	-280	802	C
ATOM	1523	O	HIS	A	212	28.433	5.403	8.847	1.00	32.84	O	
ANISOU	1523	O	HIS	A	212	3451	5298	3723	530	-300	554	O
ATOM	1524	CB	HIS	A	212	27.225	7.842	7.193	1.00	40.20	C	
ANISOU	1524	CB	HIS	A	212	4365	6313	4591	735	-333	651	C
ATOM	1525	CG	HIS	A	212	27.250	8.464	5.917	1.00	42.65	C	
ANISOU	1525	CG	HIS	A	212	4749	5665	4867	826	-350	697	C
ATOM	1526	ND1	HIS	A	212	27.092	9.812	8.925	1.00	43.91	N	

TABLE 3-continued

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ANISOU	1526	ND1	HIS	A	212	4890	6782	5032	913	-324	747	N
ATOM	1527	CD2	HIS	A	212	27.407	8.068	4.590	1.00	45.33	C	
ANISOU	1527	CD2	HIS	A	212	5110	7007	5106	838	-352	700	C
ATOM	1528	CE1	HIS	A	212	27.185	13.261	4.662	1.00	45.23	C	
ANISOU	1528	CE1	HIS	A	212	5111	5961	5115	981	-347	788	C
ATOM	1529	NE2	HIS	A	212	27.345	9.204	3.849	1.00	45.75	N	
ANISOU	1529	NE2	HIS	A	212	5196	7051	5120	9333	-389	756	N
ATOM	1530	N	SER	A	213	28.353	7.160	10.215	1.00	34.32	N	
ANISOU	1530	N	SER	A	213	3510	5447	3985	505	-217	606	N
ATOM	1531	CA	SER	A	213	28.493	6.306	11.352	1.00	34.29	C	
ANISOU	1531	CA	SER	A	213	3583	5424	4020	517	-204	570	C
ATOM	1532	C	SER	A	213	28.102	7.227	12.492	1.00	35.78	C	
ANISOU	1532	C	SER	A	213	3728	5613	4255	549	-350	582	C
ATOM	1533	O	SER	A	213	28.087	9.439	12.303	1.00	35.36	O	
ANISOU	1533	O	SER	A	213	3688	5547	4208	635	-135	814	O
ATOM	1534	CB	SER	A	213	22.976	5.835	11.457	1.00	32.35	C	
ANISOU	1534	CB	SER	A	213	3612	5272	3976	473	-170	553	C
ATOM	1535	OG	SER	A	213	30.749	6.853	12.157	1.00	32.85	O	
ANISOU	1535	OG	SER	A	213	3513	5072	3690	510	-1.00	573	O
ATOM	1536	N	PRO	A	214	27.817	5.667	13.890	1.00	35.60	N	
ANISOU	1536	N	PRO	A	214	3812	5753	4417	481	-147	556	N
ATOM	1537	CA	PRO	A	214	27.584	7.482	14.903	1.00	37.15	C	
ANISOU	1537	CA	PRO	A	214	3823	5791	4503	505	-97	559	C
ATOM	1538	C	PRO	A	214	23.712	6.511	15.156	1.00	35.29	C	
ANISOU	1538	C	PRO	A	214	3660	5457	4293	557	-39	578	C
ATOM	1539	O	PRO	A	214	28.499	9.962	15.658	1.00	33.57	O	
ANISOU	1539	O	PRO	A	214	3427	5227	4100	623	-2	591	O
ATOM	1540	CB	PRO	A	214	27.816	6.421	16.033	1.00	35.98	C	
ANISOU	1540	CB	PRO	A	214	3651	5640	4372	402	-90	529	C
ATOM	1541	CG	PRO	A	214	27.058	5.168	15.353	1.00	38.66	C	
ANISOU	1541	CG	PRO	A	214	3966	8039	4585	335	-154	512	C
ATOM	1542	CD	PRO	A	214	27.498	5.228	13.905	1.00	36.72	C	
ANISOU	1542	CD	PRO	A	214	8774	5775	4401	378	-136	523	C
ATOM	1543	N	LEU	A	215	29.535	8.033	14.873	1.00	35.02	N	
ANISOU	1543	N	LEU	A	215	3702	5347	4257	523	-28	575	N
ATOM	1544	CA	LEU	A	215	31.102	3.940	15.053	1.00	33.68	C	
ANISOU	1544	CA	LEU	A	215	3600	5083	4114	558	25	590	C
ATOM	1545	C	LEU	A	215	31.110	13.139	14.116	1.00	30.35	C	
ANISOU	1545	C	LEU	A	215	3843	5281	4311	653	34	531	C
ATOM	1546	O	LEU	A	215	31.572	11.250	14.453	1.00	37.01	O	

TABLE 3-continued

ANISOU	1546	O	LEU	A	215	4050	5431	4552	703	79	643	O
ATOM	1547	CB	LEU	A	215	32.417	8.082	15.317	100	31.71	C	
ANISOU	1547	CB	LEU	A	215	3415	4784	3889	493	33	572	C
ATOM	1548	CG	LEU	A	215	33.776	8.780	15.309	1.00	32.72	C	
ANISOU	1548	CG	LEU	A	215	3611	4794	4032	510	83	580	C
ATOM	1549	CD1	LEU	A	215	33.784	9.359	18.715	1.00	34.91	c	
ANISOU	1549	CD1	LEU	A	215	3869	5043	4354	506	126	570	C
ATOM	1550	CD2	LEU	A	215	34.978	7.768	15.217	1.00	26.46	C	
ANISOU	1550	CD2	LEU	A	215	2869	3948	3243	446	85	557	C
ATOM	1551	N	SER	A	216	30.589	9.991	12.894	1.00	39.82	N	
ANISOU	1551	N	SER	A	216	4400	5901	4320	682	-11	648	N
ATOM	1552	CA	SER	A	216	30.650	11.180	12.008	1.00	43.87	C	
ANISOU	1552	CA	SER	A	216	4951	6394	5325	775	-2	694	C
ATOM	1553	C	SER	A	216	29.895	12.392	12.508	1.00	42.67	C	
ANISOU	1553	C	SER	A	216	4761	6251	5200	853	14	713	C
ATOM	1554	O	SER	A	216	30.320	13.507	12.276	1.00	44.91	O	
ANISOU	1554	O	SER	A	216	5092	5475	5496	917	45	748	O
ATOM	1555	CB	SER	A	216	30.242	10.855	10.536	1.00	43.78	C	
ANISOU	1555	CB	SER	A	216	4889	6370	5184	799	-53	713	C
ATOM	1556	OG	SER	A	216	29.189	10.119	10.521	1.00	43.21	O	
ANISOU	1556	OG	SER	A	216	4805	6455	5155	775	-109	689	O
ATOM	1557	N	SER	A	217	28.857	12.224	13.302	1.00	40.28	N	
ANISOU	1557	N	SER	A	217	4377	6014	4915	846	0	689	N
ATOM	1558	CA	SER	A	217	28.188	13.389	13.762	1.00	40.42	C	
ANISOU	1558	CA	SER	A	217	4260	5088	4959	927	17	702	C
ATOM	1559	C	SER	A	217	28.912	13.938	15.015	1.00	44.89	C	
ANISOU	1559	C	SER	A	217	4953	5524	5580	915	81	585	C
ATOM	1560	O	SER	A	217	28.448	14.868	15.625	1.00	44.04	O	
ANISOU	1560	O	SER	A	217	4820	5412	5502	975	193	685	O
ATOM	1561	CB	SER	A	217	25.700	13.046	14.064	1.00	48.70	C	
ANISOU	1561	CB	SER	A	217	5299	7203	6002	921	-22	680	C
ATOM	1562	OG	SER	A	217	26.629	12.440	15.852	1.00	54.10	O	
ANISOU	1562	OG	SER	A	217	5941	7899	5714	857	4	539	O
ATOM	1563	N	LEU	A	218	30.072	13.989	15.397	1.00	35.24	N	
ANISOU	1563	N	LEU	A	218	2792	5247	4284	842	108	670	N
ATOM	1564	CA	LEU	A	218	30.830	13.989	15.505	1.00	27.25	C	
ANISOU	1564	CA	LEU	A	218	4081	5422	4683	837	164	655	C
ATOM	1565	C	LEU	A	218	32.225	14.287	16.041	1.00	38.53	C	
ANISOU	1565	C	LEU	A	218	4381	5486	4850	833	192	575	C
ATOM	1566	O	LEU	A	218	33.132	14.157	16.860	1.00	43.95	O	

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TABLE 3-continued

ANISOU	1566	O	LEU	A	218	5334	5099	5567	783	225	552	O
ATOM	1567	CB	LEU	A	218	80.965	12.978	17.693	1.00	28.15	C	
ANISOU	1567	CB	LEU	A	218	4154	5537	4803	744	172	610	C
ATOM	1568	CG	LEU	A	218	29.584	12.271	18.255	1.00	41.48	C	
ANISOU	1568	CG	LEU	A	218	4486	6063	5213	719	143	587	C
ATOM	1569	CD1	LEU	A	218	30.055	11.076	13.959	1.00	40.64	C	
ANISOU	1569	CD1	LEU	A	218	4377	5950	5104	612	144	557	C
ATOM	1570	CD2	LEU	A	218	29.012	13.403	19.177	1.00	42.90	C	
ANISOU	1570	CD2	LEU	A	218	4623	6257	5421	761	180	575	C
ATOM	1571	N	SER	A	219	32.380	14.638	14.755	1.00	27.47	N	
ANISOU	1571	N	SER	A	219	4223	5830	4582	872	179	715	N
ATOM	1572	CA	SER	A	219	32.718	14.172	14.117	1.00	37.09	C	
ANISOU	1572	CA	SER	A	219	4257	5231	4635	850	207	738	C
ATOM	1573	C	SER	A	219	33.810	15.124	13.465	1.00	38.91	C	
ANISOU	1573	C	SER	A	219	4533	5383	4869	942	225	789	C
ATOM	1574	O	SER	A	219	22.351	16.524	12.755	1.00	28.48	O	
ANISOU	1574	O	SER	A	219	4459	5380	4783	1009	193	819	O
ATOM	1575	CB	SER	A	219	38.921	12.708	13.023	1.00	36.22	C	
ANISOU	1575	CB	SER	A	219	4160	5131	4470	821	173	741	C
ATOM	1576	OG	SER	A	219	33.950	12.422	13.534	1.00	37.53	O	
ANISOU	1576	OG	SER	A	219	4296	5323	4537	740	154	695	O
ATOM	1577	N	THR	A	220	34.882	16.865	13.788	1.00	38.05	N	
ANISOU	1577	N	THR	A	220	4481	5175	4800	940	277	800	N
ATOM	1578	CA	THR	A	220	35.127	16.157	13.081	1.00	38.09	C	
ANISOU	1578	CA	THR	A	220	4543	5117	4817	1009	300	855	C
ATOM	1579	C	THR	A	220	35.552	17.862	11.645	1.00	39.19	C	
ANISOU	1579	C	THR	A	220	4728	5268	4594	1008	290	897	C
ATOM	1580	O	THR	A	220	35.225	18.607	10.744	1.00	35.37	O	
ANISOU	1580	O	THR	A	220	4275	4731	4382	1073	232	550	O
ATOM	1581	CB	THR	A	220	30.168	19.027	13.344	1.00	34.27	C	
ANISOU	1581	CB	THR	A	220	4135	4521	4394	995	357	851	C
ATOM	1582	OG1	THR	A	220	35.323	18.065	15.240	1.00	37.58	O	
ANISOU	1582	OG1	THR	A	220	4480	4941	4856	987	353	801	O
ATOM	1583	CG2	THR	A	220	35.225	20.498	13.393	1.00	35.05	C	
ANISOU	1583	CG2	THR	A	220	4642	4930	4896	1073	379	906	C
ATOM	1584	N	GLU	A	221	36.390	16.836	11.404	1.00	34.30	N	
ANISOU	1584	N	GLU	A	221	4197	4723	4335	934	297	872	N
ATOM	1585	CA	GLU	A	221	36.854	15.395	10.164	1.00	31.37	C	
ANISOU	1585	CA	GLU	A	221	3853	4963	3883	825	291	889	C
ATOM	1586	C	GLU	A	221	35.583	14.654	19.115	1.00	35.52	C	

TABLE 3-continued

ANISOU	1586	C	GLU	A	221	4233	4837	4325	857	258	849	C
ATOM	1587	O	GLU	A	221	95.937	14.217	11.175	1.00	33.00	O	
ANISOU	1587	O	GLU	A	221	3927	4553	4042	505	256	800	O
ATOM	1588	CB	GLU	A	221	38.316	16.349	9.940	1.00	31.94	C	
ANISOU	1588	CB	GLU	A	221	3327	4382	3325	899	553	319	C
ATOM	1589	CG	GLU	A	221	38.522	18.350	10.198	1.00	36.88	C	
ANISOU	1589	CG	GLU	A	221	4554	4813	4501	943	357	901	C
ATOM	1590	CD	GLU	A	221	38.135	19.358	9.121	45.37	C		
ANISOU	1590	CD	GLU	A	221	5840	6014	5753	1322	391	1036	C
ATOM	1591	OE1	GLU	A	221	37.583	18.901	8.071	1.00	41.04	O	
ANISOU	1591	OE1	GLU	A	221	5156	5414	5015	1046	352	1058	O
ATOM	1592	OE2	GLU	A	221	38.351	20.632	9.327	1.00	47.32	O	
ANISOU	1592	OE2	GLU	A	221	5076	6127	6003	1060	425	1073	O
ATOM	1593	N	MET	A	222	36.941	34.252	8.920	1.00	32.33	N	
ANISOU	1593	N	MET	A	222	3901	4523	3555	555	235	551	N
ATOM	1594	CA	MET	A	222	37.053	12.787	8.858	1.00	40.53	C	
ANISOU	1594	CA	MET	A	222	4838	5554	4840	790	203	810	C
ATOM	1595	C	MET	A	222	38.004	12.428	7.724	1.00	40.55	C	
ANISOU	1595	C	MET	A	222	4975	5607	4524	777	222	820	C
ATOM	1596	O	MET	A	222	37.985	13.079	6.705	1.00	41.53	O	
ANISOU	1596	O	MET	A	222	5975	5573	4839	826	230	870	O
ATOM	1597	CB	MET	A	222	55.108	12.167	8.430	1.00	44.61	C	
ANISOU	1597	CB	MET	A	222	5385	6223	5335	805	133	800	C
ATOM	1598	CG	MET	A	222	34.771	11.743	9.544	1.00	54.18	C	
ANISOU	1598	CG	MET	A	222	5524	7480	5582	780	132	763	C
ATOM	1599	SD	MET	A	222	33.983	15.126	9.175	1.00	54.33	S	
ANISOU	1599	SD	MET	A	222	6560	7655	6617	724	23	716	S
ATOM	1600	CE	MET	A	222	33.384	9.717	10.821	1.00	49.94	C	
ANISOU	1600	CE	MET	A	222	5368	7048	6058	673	25	575	C
ATOM	1601	N	LEU	A	223	35.793	11.353	7687	1.00	37.07	N	
ANISOU	1601	N	LEU	A	223	4514	5231	4466	715	227	773	N
ATOM	1602	CA	LEU	A	223	59.557	39.432	6.731	1.00	43.48	C	
ANISOU	1602	CA	LEU	A	223	5013	5600	4757	704	235	771	C
ATOM	1603	C	LEU	A	223	39.032	9.402	6.735	1.00	38.74	C	
ANISOU	1603	C	LEU	A	223	4761	5439	4520	663	174	715	C
ATOM	1604	O	LEU	A	223	38.871	5.525	7.743	1.00	34.30	O	
ANISOU	1604	O	LEU	A	223	4151	4865	4004	620	158	678	O
ATOM	1605	CB	LEU	A	223	41.038	10.885	6.934	1.30	42.92	C	
ANISOU	1605	CB	LEU	A	223	5955	5837	5115	670	239	769	C
ATOM	1606	CG	LEU	A	223	41.712	12.255	6.8929	1.00	45.92	C	

TABLE 3-continued

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ANISOU	1606	CG	LEU	A	223	5774	6152	5321	703	364	835	C
ATOM	1607	CD1	LEU	A	223	43.221	12.124	6.847	1.00	38.40	C	
ANISOU	1607	CD1	LEU	A	223	4846	5146	4000	660	423	799	C
ATOM	1608	CD2	LEU	A	223	41.239	13.039	5.599	1.00	47.31	C	
ANISOU	1608	CD2	LEU	A	223	5939	6358	5629	762	365	882	C
ATOM	1609	N	VAL	A	224	38.692	8.913	5.520	1.00	35.69	N	
ANISOU	1609	N	VAL	A	224	4391	5112	4056	678	139	715	N
ATOM	1610	CA	VAL	A	224	38.053	7.619	5.337	1.00	33.42	C	
ANISOU	1610	CA	VAL	A	224	4077	4885	3737	647	72	664	C
ATOM	1611	C	VAL	A	224	38.975	6.699	4.534	1.00	34.94	C	
ANISOU	1611	C	VAL	A	224	4310	5077	3839	617	75	523	C
ATOM	1612	O	VAL	A	224	39.263	6.987	3.417	1.00	34.78	O	
ANISOU	1612	O	VAL	A	224	4333	5075	3806	652	93	652	O
ATOM	1613	CB	VAL	A	224	35.672	7.326	4.355	1.00	33.13	C	
ANISOU	1613	CB	VAL	A	224	4649	5554	4275	688	12	589	C
ATOM	1614	CG1	VAL	A	224	36.035	6.473	4.247	1.00	34.99	C	
ANISOU	1614	CG1	VAL	A	224	4227	5231	3836	648	-62	635	C
ATOM	1615	CG2	VAL	A	224	35.783	8.655	5.593	1.00	32.84	C	
ANISOU	1615	CG2	VAL	A	224	7923	4895	3656	713	3	715	C
ATOM	1616	N	ALA	A	225	34.314	5.572	5.595	1.00	33.01	N	
ANISOU	1616	N	ALA	A	225	4353	4813	3575	559	60	559	N
ATOM	1617	CA	ALA	A	225	45.116	4.545	4.365	1.00	35.35	C	
ANISOU	1617	CA	ALA	A	225	4385	5111	3939	537	57	521	C
ATOM	1618	C	ALA	A	225	39.341	3.552	3.555	1.00	37.67	C	
ANISOU	1618	C	ALA	A	225	4668	5474	4170	523	-20	486	C
ATOM	1619	O	ALA	A	225	37.993	3.399	4.170	1.00	39.05	O	
ANISOU	1619	O	ALA	A	225	4795	5684	4351	511	-75	435	O
ATOM	1620	CB	ALA	A	225	40.985	3.791	5.454	1.00	32.85	C	
ANISOU	1620	CB	ALA	A	225	4064	4772	3701	480	71	473	C
ATOM	1621	N	ALA	A	226	34.581	2.870	2.619	1.00	35.81	N	
ANISOU	1621	N	ALA	A	226	4473	5261	3873	529	-33	453	N
ATOM	1622	CA	ALA	A	226	78.600	1.574	1.945	1.00	39.341	C	
ANISOU	1622	CA	ALA	A	226	4917	5767	4263	514	-109	210	C
ATOM	1623	C	ALA	A	226	33.612	0.632	2.716	1.00	33.54	C	
ANISOU	1623	C	ALA	A	226	4155	5310	3575	446	-157	350	C
ATOM	1624	O	ALA	A	226	79.210	0.466	3.841	1.00	76.40	O	
ANISOU	1624	O	ALA	A	226	4508	5758	4015	412	-129	338	O
ATOM	1625	CB	ALA	A	226	39.422	1.528	0.581	1.00	38.22	C	
ANISOU	1625	CB	ALA	A	226	4872	5452	4073	543	-101	787	C
ATOM	1626	N	ARG	A	227	37.720	-0.235	2.317	1.00	33.51	N	

TABLE 3-continued

ANISOU 1626 N ARG A 227	4140 5053 3576 423 -233 313 N
ATOM 1627 CA ARG A 227	37.369 -1.523 3.010 1.00 34.30 C
ANISOU 1627 CA ARG A 227	4217 5109 3678 251 -293 256 C
ATOM 1628 C ARG A 227	35.742 -1.367 4.381 1.00 33.88 C
ANISOU 1628 C ARG A 227	4107 5355 9597 311 -295 277 C
ATOM 1629 O ARG A 227	37.409 -1.646 5.374 1.00 33.47 O
ANISOU 1629 O ARG A 227	4057 4551 3710 277 -268 265 O
ATOM 1630 CB ARG A 227	38.560 -2.417 3.227 1.00 37.30 C
ANISOU 1630 CB ARG A 227	4750 5510 4132 322 -270 203 C
ATOM 1631 CG ARG A 227	39.453 -2.457 2.011 1.00 44.09 C
ANISOU 1631 CG ARG A 227	5552 3314 4885 365 -243 182 C
ATOM 1632 CD ARG A 227	39.299 -3.775 1.245 1.00 58.42 C
ANISOU 1632 CD ARG A 227	7396 8147 6655 342 -308 108 C
ATOM 1633 NE ARG A 227	38.505 -3.532 0.041 1.00 53.27 N
ANISOU 1633 NE ARG A 227	5751 7580 6539 377 -345 117 N
ATOM 1634 CZ ARG A 227	38.889 -2.621 -1.040 1.00 59.64 C
ANISOU 1634 CZ ARG A 227	7545 8326 6540 439 -336 142 C
ATOM 1635 NH1 ARG A 227	40.372 -2.244 -1.159 1.00 52.61 N
ANISOU 1635 NH1 ARG A 227	5704 7501 5753 472 -222 163 N
ATOM 1636 NH2 ARG A 227	33.044 -2.694 -2.040 1.00 61.05 N
ANISOU 1636 NH2 ARG A 227	7779 8683 6735 466 -353 149 N
ATOM 1637 N PRO A 228	35.410 -1.099 4.419 1.00 35.30 N
ANISOU 1637 N PRO A 228	4234 5315 3863 311 -339 299 N
ATOM 1638 CA PRO A 228	34.640 -1.078 5.655 1.00 34.16 C
ANISOU 1638 CA PRO A 228	4028 5172 3779 268 -350 312 C
ATOM 1639 C PRO A 228	34.776 -2.426 6.330 1.00 34.46 C
ANISOU 1639 C PRO A 228	4068 5168 3857 185 -384 261 C
ATOM 1640 O PRO A 228	34.767 -3.470 5.675 1.00 34.02 O
ANISOU 1640 O PRO A 228	4038 5116 3772 156 -434 212 O
ATOM 1641 CB PRO A 228	33.204 -0.977 5.149 1.00 36.75 C
ANISOU 1641 CB PRO A 228	4304 5594 4066 277 -411 322 C
ATOM 1642 CG PRO A 228	33.251 -0.381 3.808 1.00 34.24 C
ANISOU 1642 CG PRO A 228	4019 5319 3672 348 -413 340 C
ATOM 1643 CD PRO A 228	34.598 -0.714 3.237 1.00 34.61 C
ANISOU 1643 CD PRO A 228	4141 5311 3698 358 -377 315 C
ATOM 1644 N GLU A 229	34.846 -2.455 7.652 1.00 31.28 N
ANISOU 1644 N GLU A 229	3641 4723 3520 145 -362 271 N
ATOM 1645 CA GLU A 229	34.897 -3.779 8.326 1.00 29.28 C
ANISOU 1645 CA GLU A 229	3394 4428 3304 62 -399 229 C
ATOM 1646 C GLU A 229	33.505 -4.339 8.517 1.00 30.23 C

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TABLE 3-continued

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ANISOU	1646	C	GLU	A	229	3457	4608	3422	5	-461	222	C
ATOM	1647	O	GLU	A	229	32.571	-3.569	8.609	1.00	35.50	O	
ANISOU	1647	O	GLU	A	229	4066	5341	4081	27	-460	255	O
ATOM	1648	CB	GLU	A	229	35.556	-3.591	9.734	1.00	30.45	C	
ANISOU	1648	CB	GLU	A	229	3542	4507	3520	39	-351	246	C
ATOM	1649	CG	GLU	A	229	37.015	-3.264	9.509	1.00	30.55	C	
ANISOU	1649	CG	GLU	A	229	3611	4457	3541	84	-299	241	C
ATOM	1650	CD	GLU	A	229	37.834	-3.356	10.735	1.00	33.82	C	
ANISOU	1650	CD	GLU	A	229	4036	4797	4017	58	-266	242	C
ATOM	1651	OE1	GLU	A	229	37.289	-3.191	11.856	1.00	35.26	O	
ANISOU	1651	OE1	GLU	A	229	4181	4983	4234	234	-262	264	O
ATOM	1652	OE2	GLU	A	229	39.050	-3.531	10.600	1.00	33.05	O	
ANISOU	1652	OE2	GLU	A	229	3983	4642	3934	75	-241	222	O
ATOM	1653	N	GLY	A	230	33.358	-5.660	8.624	1.00	28.87	N	
ANISOU	1653	N	GLY	A	230	3298	4412	3261	-70	-515	178	N
ATOM	1654	CA	GLY	A	230	32.047	-6.313	8.917	1.00	31.12	C	
ANISOU	1654	CA	GLY	A	230	3524	4748	3551	-143	-574	170	C
ATOM	1655	C	GLY	A	230	32.266	-7.090	10.214	1.00	34.23	C	
ANISOU	1655	C	GLY	A	230	2925	5076	4005	-223	-570	167	C
ATOM	1656	O	GLY	A	230	33.407	-7.113	10.727	1.00	36.84	O	
ANISOU	1656	O	GLY	A	230	4305	5326	4365	-212	-531	166	O
ATOM	1657	N	PRO	A	231	31.216	-7.768	10.716	1.00	33.86	N	
ANISOU	1657	N	PRO	A	231	3833	5063	3978	-305	-613	163	N
ATOM	1658	CA	PRO	A	231	31.263	-8.417	11.993	1.00	34.91	C	
ANISOU	1658	CA	PRO	A	231	3965	5142	4159	-384	-608	172	C
ATOM	1659	C	PRO	A	231	32.373	-9.455	12.062	1.00	35.15	C	
ANISOU	1659	C	PRO	A	231	4080	5066	4210	-412	-625	136	C
ATOM	1660	O	PRO	A	231	32.851	-9.742	13.161	1.00	29.72	O	
ANISOU	1660	O	PRO	A	231	3414	4316	3564	-448	-604	151	O
ATOM	1661	CB	PRO	A	231	29.869	-9.107	12.082	1.00	34.13	C	
ANISOU	1661	CB	PRO	A	231	3802	5105	4060	-471	-665	165	C
ATOM	1662	CG	PRO	A	231	28.957	-8.128	11.803	1.00	36.11	C	
ANISOU	1662	CG	PRO	A	231	3984	5466	4270	-410	-668	179	C
ATOM	1663	CD	PRO	A	231	29.879	-7.927	10.087	1.00	37.15	C	
ANISOU	1663	CD	PRO	A	231	4184	5571	4362	-330	-670	154	C
ATOM	1664	N	LEU	A	232	32.713	-10.073	10.934	1.00	33.04	N	
ANISOU	1664	N	LEU	A	232	3860	4781	3914	-397	-667	88	N
ATOM	1665	CA	LEU	A	232	33.667	-11.165	10.939	1.00	35.50	C	
ANISOU	1665	CA	LEU	A	232	4249	4993	4247	-423	-692	47	C
ATOM	1666	C	LEU	A	232	35.025	-10.741	10.308	1.00	36.14	C	

TABLE 3-continued

ANISOU	1666	C	LEU	A	232	4386	5032	4313	-333	-652	28	C
ATOM	1667	O	LEU	A	232	35.934	-11.587	10.206	1.00	34.68	O	
ANISOU	1667	O	LEU	A	232	4265	4766	4144	-337	-669	-13	O
ATOM	1668	CB	LEU	A	232	33.128	-12.341	10.122	1.00	36.97	C	
ANISOU	1668	CB	LEU	A	232	4455	5179	4414	-477	-774	-7	C
ATOM	1669	CG	LEU	A	232	31.975	-13.121	10.744	1.00	40.58	C	
ANISOU	1669	CG	LEU	A	232	4870	5651	4896	-587	-823	0	C
ATOM	1670	CD1	LEU	A	232	31.421	-14.171	9.766	1.00	38.22	C	
ANISOU	1670	CD1	LEU	A	232	4589	5358	4574	-634	-906	-59	C
ATOM	1671	CD2	LEU	A	232	32.517	-13.761	12.019	1.00	38.34	C	
ANISOU	1671	CD2	LEU	A	232	4624	5274	4668	-645	-812	17	C
ATOM	1672	N	THR	A	233	35.131	-9.486	9.839	1.00	30.76	N	
ANISOU	1672	N	THR	A	233	3682	4406	3601	-254	-602	56	N
ATOM	1673	CA	THR	A	233	36.424	-8.979	9.342	1.00	34.57	C	
ANISOU	1673	CA	THR	A	233	4210	4852	4072	-175	-552	47	C
ATOM	1674	C	THR	A	233	37.028	-7.809	10.138	1.00	33.17	C	
ANISOU	1674	C	THR	A	233	4017	4661	3924	-131	-476	96	C
ATOM	1675	O	THR	A	233	37.707	-6.950	9.615	1.00	34.74	O	
ANISOU	1675	O	THR	A	233	4228	4866	4104	-61	-427	108	O
ATOM	1676	CB	THR	A	233	36.223	-8.649	7.821	1.00	37.93	C	
ANISOU	1676	CB	THR	A	233	4643	5343	4427	-117	-564	28	C
ATOM	1677	OG1	THR	A	233	35.202	-7.675	7.736	1.00	37.25	O	
ANISOU	1677	OB1	THR	A	233	4495	5343	4316	-98	-556	74	O
ATOM	1678	CG2	THR	A	233	35.715	-9.893	7.021	1.00	40.75	C	
ANISOU	1678	CG2	THR	A	233	5023	5707	4755	-162	-646	-32	C
ATOM	1679	N	GLY	A	234	36.865	-7.763	11.472	1.00	29.36	N	
ANISOU	1679	N	GLY	A	234	3511	4154	3489	-175	-462	126	N
ATOM	1680	CA	GLY	A	234	37.455	-6.636	12.131	1.00	32.28	C	
ANISOU	1680	CA	GLY	A	234	3870	4511	3883	-130	-393	165	C
ATOM	1681	C	GLY	A	234	38.955	-6.871	12.092	1.00	31.45	C	
ANISOU	1681	C	GLY	A	234	3821	4328	3801	-101	-369	138	C
ATOM	1682	O	GLY	A	234	39.401	-8.018	12.102	1.00	29.10	O	
ANISOU	1682	O	GLY	A	234	3565	3974	3519	-133	-408	98	O
ATOM	1683	N	GLY	A	235	39.721	-5.803	12.038	1.00	31.17	N	
ANISOU	1683	N	GLY	A	235	3787	4287	3770	-39	-307	159	N
ATOM	1684	CA	GLY	A	235	41.177	-5.998	12.087	1.00	34.52	C	
ANISOU	1684	CA	GLY	A	235	4254	4640	4222	-14	-281	134	C
ATOM	1685	C	GLY	A	235	41.869	-6.459	10.827	1.00	35.18	C	
ANISOU	1685	C	GLY	A	235	4377	4716	4273	22	-287	89	C
ATOM	1686	O	GLY	A	235	43.098	-6.501	10.860	1.00	32.01	O	

TABLE 3-continued

ANISOU	1686	O	GLY	A	235	4001	4264	3897	50	-258	68	O
ATOM	1687	N	ALA	A	236	41.115	-6.776	9.729	1.00	29.12	N	
ANISOU	1687	N	ALA	A	236	3612	4003	3450	25	-324	71	N
ATOM	1688	CA	ALA	A	236	41.665	-7.259	8.493	1.00	33.61	C	
ANISOU	1688	CA	ALA	A	236	4220	4573	3976	59	-333	23	C
ATOM	1689	C	ALA	A	236	42.781	-6.321	6.037	1.00	32.76	C	
ANISOU	1689	C	ALA	A	236	4124	4463	3862	126	-258	35	C
ATOM	1690	O	ALA	A	236	42.672	-5.072	8.100	1.00	34.03	O	
ANISOU	1690	O	ALA	A	236	4258	4654	4016	158	-207	87	O
ATOM	1691	CB	ALA	A	236	40.580	-7.328	7.352	1.00	33.65	C	
ANISOU	1691	CB	ALA	A	236	4219	4657	3910	63	-374	15	C
ATOM	1692	N	PHE	A	237	43.844	-6.905	7.510	1.00	29.62	N	
ANISOU	1692	N	PHE	A	237	3764	4029	3463	150	-250	-15	N
ATOM	1693	CA	PHE	A	237	44-978	4029	3463	150	-250	-15	C
ANISOU	1693	CA	PHE	A	237	3842	4098	3538	205	-173	-4	C
ATOM	1694	C	PHE	A	237	44.599	-5.101	6.007	1.00	32.94	C	
ANISOU	1694	C	PHE	A	237	4186	4520	3809	251	-140	28	C
ATOM	1695	O	PHE	A	237	45.025	-3.984	4.965	1.00	34.81	O	
ANISOU	1695	O	PHE	A	237	4413	4767	4048	285	-75	71	O
ATOM	1696	CB	PHE	A	237	46.166	-6.952	6.851	1.00	33.29	C	
ANISOU	1686	CB	PHE	A	237	4268	4439	3942	222	-173	-70	C
ATOM	1697	CG	PHE	A	237	47.398	-6.237	6.393	1.00	31.86	C	
ANISOU	1697	CG	PHE	A	237	4087	4253	3760	275	-95	-68	C
ATOM	1698	CD1	PHE	A	237	47.953	-5.167	7.116	1.00	31.99	C	
ANISOU	1698	CD1	PHE	A	237	4075	4257	3823	284	-34	-21	C
ATOM	1699	CD2	PHE	A	237	48.016	-5.628	5.175	1.00	31.38	C	
ANISOU	1699	CD2	PHE	A	237	4055	4219	3849	315	-78	-118	C
ATOM	1700	CE1	PHE	A	237	49.135	-4.549	8.698	1.00	32.26	C	
ANISOU	1700	CE1	PHE	A	237	4107	4290	3861	325	39	-21	C
ATOM	1701	CE2	PHE	A	237	49.196	-5.998	4.775	1.00	31.52	C	
ANISOU	1701	CE2	PHE	A	237	4082	4252	3681	369	0	-117	C
ATOM	1702	CZ	PHE	A	237	49.737	-4.974	5.434	1.00	35.02	C	
ANISOU	1702	CZ	PHE	A	237	4481	4663	4161	361	58	-68	C
ATOM	1703	N	ALA	A	238	43.733	-5.532	5.172	1.00	32.19	N	
ANISOU	1703	N	ALA	A	238	4103	4477	3651	249	-188	10	N
ATOM	1704	CA	ALA	A	238	43.354	-4.858	4.048	1.00	40.10	C	
ANISOU	1704	CA	ALA	A	238	5107	5551	4579	295	-163	44	C
ATOM	1705	C	ALA	A	238	42.563	-3.397	4.575	1.00	40.82	C	
ANISOU	1705	C	ALA	A	238	5157	5672	4681	301	-141	121	C
ATOM	1706	O	ALA	A	238	42.590	-2.305	4.001	1.00	36.66	O	

TABLE 3-continued

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ANISOU	1706	O	ALA	A	238	4634	5177	4119	347	-91	167	O
ATOM	1707	CB	ALA	A	238	42.609	-5.428	2.937	1.00	41.88	C	
ANISOU	1707	CB	ALA	A	238	5329	5803	4703	294	-226	2	C
ATOM	1708	N	SER	A	239	41.853	-3.522	5.594	1.00	35.52	N	
ANISOU	1708	N	SER	A	239	4451	4985	4050	256	-172	136	N
ATOM	1709	CA	SER	A	239	41.206	-2.357	6.314	1.00	34.89	C	
ANISOU	1709	CA	SER	A	239	4331	4927	3999	265	-147	201	C
ATOM	1710	C	SER	A	239	42.202	-1.473	7.092	1.00	32.55	C	
ANISOU	1710	C	SER	A	239	4045	4583	3773	282	-75	231	C
ATOM	1711	O	SER	A	239	42.158	-0.232	6.989	1.00	31.59	O	
ANISOU	1711	O	SER	A	239	3901	4970	3631	321	-28	284	O
ATOM	1712	CB	SER	A	239	40.110	-2.795	7.252	1.00	33.93	C	
ANISOU	1712	CB	SER	A	239	4170	4814	3909	211	-200	204	C
ATOM	1713	OG	SER	A	239	39.135	-3.426	6.455	1.00	31.01	O	
ANISOU	1713	OG	SER	A	239	3796	4502	3488	198	-264	182	O
ATOM	1714	N	LYS	A	240	43.123	-2.092	7.800	1.00	30.82	N	
ANISOU	1714	N	LYS	A	240	3823	4292	3595	256	-68	198	N
ATOM	1715	CA	LYS	A	240	44.061	-1.340	8.597	1.00	31.64	C	
ANISOU	1715	CA	LYS	A	240	3920	4345	3755	256	-9	220	C
ATOM	1716	C	LYS	A	240	45.132	-0.667	7.729	1.00	31.46	C	
ANISOU	1716	C	LYS	A	240	3923	4320	3713	313	57	227	C
ATOM	1717	O	LYS	A	240	45.846	0.386	8.072	1.00	30.20	O	
ANISOU	1717	O	LYS	A	240	3753	4139	3584	331	114	264	O
ATOM	1718	CB	LYS	A	240	44.760	-2.259	9.611	1.00	28.42	C	
ANISOU	1718	CB	LYS	A	240	3518	3871	2411	225	-28	180	C
ATOM	1719	CG	LYS	A	240	43.830	-2.930	10.661	1.00	32.44	C	
ANISOU	1719	CG	LYS	A	240	4008	4373	3946	168	-86	178	C
ATOM	1720	CD	LYS	A	240	43.003	-1.915	11.529	1.00	29.16	C	
ANISOU	1720	CD	LYS	A	240	3553	3580	3545	163	-69	231	C
ATOM	1721	CE	LYS	A	240	41.557	-1.303	11.071	1.00	35.50	C	
ANISOU	1721	CE	LYS	A	240	4330	4856	4304	160	-103	252	C
ATOM	1722	NZ	LYS	A	240	40.721	-0.389	12.083	1.00	29.77	N	
ANISOU	1722	NZ	LYS	A	240	3561	4150	3602	151	-90	794	N
ATOM	1723	N	VAL	A	241	45.463	-1.273	6.592	1.00	37.05	N	
ANISOU	1723	N	VAL	A	241	4651	5050	4360	331	51	191	N
ATOM	1724	CA	VAL	A	241	46.545	-0.720	5.799	1.00	33.73	C	
ANISOU	1724	CA	VAL	A	241	4637	5009	4309	371	119	195	C
ATOM	1725	C	VAL	A	241	46.115	7.624	5.252	1.00	35.06	C	
ANISOU	1725	C	VAL	A	241	4425	4835	4055	406	161	264	C
ATOM	1726	O	VAL	A	241	46.909	1.558	5.197	1.00	35.63	O	

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ANISOU	1726	O	VAL	A	241	4500	4890	4146	425	230	297	O
ATOM	1727	CB	VAL	A	241	47.059	-1.717	4.730	1.00	37.45	C	
ANISOU	1727	CB	VAL	A	241	4761	5338	4351	385	108	304	C
ATOM	1728	CG1	VAL	A	241	46.159	-1.724	3.532	1.00	73.85	C	
ANISOU	1728	CG1	VAL	A	241	4326	4775	3371	408	80	142	C
ATOM	1729	CG2	VAL	A	241	48.492	-1.302	4.318	1.00	37.55	C	
ANISOU	1729	CG2	VAL	A	241	4795	7125	4385	414	187	123	C
ATOM	1730	N	GLY	A	242	44.857	0.783	4.890	1.00	32.25	N	
ANISOU	1730	N	GLY	A	242	4065	4575	3657	414	119	290	N
ATOM	1731	CA	GLY	A	242	44.480	2.114	4.418	1.00	32.41	C	
ANISOU	1731	CA	GLY	A	242	4085	4535	3540	454	156	350	C
ATOM	1732	C	GLY	A	242	44.390	3.155	5.516	1.00	35.41	C	
ANISOU	1732	C	GLY	A	242	4443	4926	4086	451	187	406	C
ATOM	1733	O	GLY	A	242	44.645	4.340	8.258	1.00	36.59	O	
ANISOU	1733	O	GLY	A	242	4501	5069	4232	483	241	457	O
ATOM	1734	N	ALA	A	243	43.991	2.713	6.720	1.00	23.72	N	
ANISOU	1734	N	ALA	A	243	3562	4054	3297	412	150	387	N
ATOM	1735	CA	ALA	A	243	43.941	3.625	7.947	1.00	32.55	C	
ANISOU	1735	CA	ALA	A	243	4059	4537	3888	406	177	421	C
ATOM	1736	C	ALA	A	243	45.379	4.108	8.247	1.00	32.57	C	
ANISOU	1736	C	ALA	A	243	4074	4479	3902	406	244	420	C
ATOM	1737	O	ALA	A	243	45.631	5.325	8.433	1.00	28.59	O	
ANISOU	1737	O	ALA	A	243	3527	3906	3417	427	294	455	O
ATOM	1738	CB	ALA	A	243	43.298	2.942	3.142	1.00	20.96	C	
ANISOU	1738	CB	ALA	A	243	3280	3782	3183	361	127	395	C
ATOM	1739	N	LEU	A	244	46.346	3.390	8.136	1.00	31.93	N	
ANISOU	1739	N	LEU	A	244	3927	4297	3794	388	245	370	N
ATOM	1740	CA	LEU	A	244	47.747	3.891	8.310	1.00	31.90	C	
ANISOU	1740	CA	LEU	A	244	3955	4282	3875	389	303	355	C
ATOM	1741	C	LEU	A	244	43.274	4.524	7.225	1.00	32.18	C	
ANISOU	1741	C	LEU	A	244	4022	4332	3672	424	374	403	C
ATOM	1742	O	LEU	A	244	49.725	5.437	7.551	1.00	32.69	O	
ANISOU	1742	O	LEU	A	244	4082	4351	3981	425	431	429	O
ATOM	1743	CB	LEU	A	244	48.561	2.362	8.443	1.00	70.38	C	
ANISOU	1743	CB	LEU	A	244	3334	4123	3766	367	290	296	C
ATOM	1744	CG	LEU	A	244	48.337	1.492	9.053	1.00	30.57	C	
ANISOU	1744	CG	LEU	A	244	3781	4055	3778	327	233	264	C
ATOM	1745	CD1	LEU	A	244	49.054	0.055	9.419	1.00	32.12	C	
ANISOU	1745	CD1	LEU	A	244	3092	4236	9976	316	199	196	C
ATOM	1746	CD2	LEU	A	244	43.804	2.126	11.015	1.00	28.32	C	

TABLE 3-continued

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ANISOU 1746 CD2 LEO A 244	3448 3593 3543 308 252 279	C
ATOM 1747 N LEU A 245	47.940 4.272 5.942 1.00 31.23	N
ANISOU 1747 N LEU A 245	3929 4258 3570 450 357 406	N
ATOM 1748 CA LEU A 245	48.239 5.214 4.455 1.00 35.78	C
ANISOU 1748 CA LEU A 245	4522 4356 4195 434 427 456	C
ATOM 1749 C LEU A 245	47.709 6.860 5.157 1.00 34.98	C
ANISOU 1749 C LEU A 245	4432 4750 4110 503 452 531	C
ATOM 1750 O LEU A 245	48.435 7.629 4.989 1.00 33.78	O
ANISOU 1750 O LEU A 245	4292 4568 3975 511 510 570	O
ATOM 1751 CB LEU A 245	47.522 4.725 7.581 1.00 31.35	C
ANISOU 1751 CB LEU A 245	4001 4378 8535 511 397 451	C
ATOM 1752 CG LEU A 245	47.756 5.667 2.372 1.00 39.57	C
ANISOU 1752 CG LEU A 245	5079 5450 4505 549 451 509	C
ATOM 1753 CD1 LEU A 245	49.205 5.782 1.958 1.00 47.82	C
ANISOU 1753 CD1 LEU A 245	5243 5539 4572 545 531 499	C
ATOM 1754 CD2 LEU A 245	46.925 5.164 1.157 1.00 34.43	C
ANISOU 1754 CD2 LEU A 245	4455 4675 3749 577 405 504	C
ATOM 1755 N LEU A 246	46.456 6.783 5.602 1.00 36.38	N
ANISOU 1755 N LEU A 246	4554 4544 4284 510 400 548	N
ATOM 1756 CA LEU A 246	45.960 8.103 6.099 1.00 40.90	C
ANISOU 1756 CA LEU A 246	5161 5492 4686 530 419 608	C
ATOM 1757 C LEU A 246	46.681 8.747 7.115 1.00 38.74	C
ANISOU 1757 C LEU A 246	4875 5144 4699 507 459 610	C
ATOM 1758 O LEU A 246	47.125 9.975 7.013 1.00 56.75	O
ANISOU 1758 O LEU A 246	4640 4860 4403 527 515 563	O
ATOM 1759 CB LEU A 246	44.571 8.011 5.751 1.00 40.07	C
ANISOU 1759 CB LEU A 246	5925 5413 4797 553 365 610	C
ATOM 1760 CG LEU A 246	6370 7.332 6.032 1.00 50.72	C
ANISOU 1760 CG LEU A 246	5370 5358 5063 543 289 603	C
ATOM 1761 CD1 LEU A 246	42.171 7.371 7.013 1.00 56.72	C
ANISOU 1761 CD1 LEU A 246	7086 7014 6851 542 239 553	C
ATOM 1762 CD2 LEU A 246	43.085 7 965 4.700 1.00 47.40	C
ANISOU 1762 CD2 LEU A 246	5985 6452 5551 595 295 652	C
ATOM 1763 N VAL A 247	47.331 7.373 8.118 1.00 33.57	N
ANISOU 1763 N VAL A 247	4194 4460 4102 463 449 555	N
ATOM 1764 CA VAL A 247	48.305 8.451 9.110 1.00 34.16	C
ANISOU 7 1764 CA VAL A 247	4254 4455 4259 443 490 549	C
ATOM 1765 C VAL A 247	49.583 8.950 6.400 1.00 34.23	C
ANISOU 1765 C VAL A 247	4281 4453 4273 444 551 562	C
ATOM 1766 O VAL A 247	50.143 10.097 8.659 1.00 51.52	O

TABLE 3-continued

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ANISOU	1766	O	VAL	A	247	3941	4060	3974	442	615	597	O
ATOM	1767	CB	VAL	A	247	48.694	7.405	10.218	1.00	30.44	C	
ANISOU	1767	CB	VAL	A	247	3755	3969	3641	402	454	487	C
ATOM	1768	CG1	VAL	A	247	49.348	7.986	11.039	1.00	30.37	C	
ANISOU	1768	CG1	VAL	A	247	3734	3896	3909	380	456	477	C
ATOM	1769	CG2	VAL	A	247	47.533	7.184	11.209	1.00	23.55	C	
ANISOU	1769	CG2	VAL	A	247	3496	3742	3514	390	395	430	C
ATOM	1770	N	ASP	A	248	50.076	8.121	7.476	1.00	32.34	N	
ANISOU	1770	N	ASP	A	248	4052	4249	3985	445	556	533	N
ATOM	1771	CA	ASP	A	248	51.334	8.493	5.809	1.00	31.81	C	
ANISOU	1771	CA	ASP	A	248	3935	4159	3924	443	640	540	C
ATOM	1772	C	ASP	A	248	51.224	9.764	5.992	1.00	35.32	C	
ANISOU	1772	C	ASP	A	248	4473	4616	4333	468	695	616	C
ATOM	1773	O	ASP	A	248	52.107	10.838	6.108	1.00	31.84	O	
ANISOU	1773	O	ASP	A	248	4032	4130	3937	453	759	643	O
ATOM	1774	CB	ASP	A	248	51.814	7.333	5.919	1.00	35.37	C	
ANISOU	1774	CB	ASP	A	248	4451	4664	4323	443	630	490	C
ATOM	1775	CG	ASP	A	248	53.201	7.544	5.397	1.00	37.81	C	
ANISOU	1775	CG	ASP	A	248	4757	4964	4643	439	713	484	C
ATOM	1776	OD1	ASP	A	248	54.190	7.875	5.145	1.00	34.80	O	
ANISOU	1776	OD1	ASP	A	248	4345	4532	4344	412	749	471	O
ATOM	1777	OD2	ASP	A	248	53.205	7.248	4.255	1.00	32.43	O	
ANISOU	1777	OD2	ASP	A	248	4999	4332	3592	459	733	453	O
ATOM	1778	N	VAL	A	249	55.165	9.344	5.160	1.00	35.34	N	
ANISOU	1778	N	VAL	A	249	4314	4475	4064	503	667	652	N
ATOM	1779	CA	VAL	A	249	49.830	11.026	4.359	1.00	37.23	C	
ANISOU	1779	CA	VAL	A	249	4786	4916	4454	534	706	731	C
ATOM	1780	C	VAL	A	249	49.638	12.262	5.225	1.00	38.13	C	
ANISOU	1780	C	VAL	A	249	4895	4963	4631	536	719	777	C
ATOM	1781	O	VAL	A	249	50.165	13.546	4.941	1.00	41.42	O	
ANISOU	1781	O	VAL	A	249	5537	5340	5062	537	751	351	O
ATOM	1782	CB	VAL	A	249	48.653	10.756	3.406	1.00	39.70	C	
ANISOU	1782	CB	VAL	A	249	5122	5297	4665	575	655	753	C
ATCOM	1783	CG1	VAL	A	249	48.385	12.001	2.632	1.00	44.42	C	
ANISOU	1783	CG1	VAL	A	249	5765	5895	5219	512	890	840	C
ATOM	1784	CG2	VAL	A	249	44.995	9	589	2.446	1.00	35.63	C
ANISOU	1784	CG2	VAL	A	249	4749	4976	4210	576	646	705	C
ATOM	1785	N	LEU	A	250	48.994	12.092	6.372	1.00	38.89	N	
ANISOU	1785	N	LEU	A	250	4959	5040	4776	529	669	750	N
ATOM	1786	CA	LEU	A	250	48.843	13.227	7.246	1.00	33.59	C	

TABLE 3-continued

ANISOU	1786	CA	LEU	A	250	4287	4307	4170	532	683	781	C
ATOM	1787	C	LEU	A	250	50.197	13.773	7.623	1.00	32.20	C	
ANISOU	1787	C	LEU	A	250	4107	4065	4063	495	749	779	C
ATOM	1788	O	LEU	A	250	50.416	14.949	7.514	1.00	30.97	O	
ANISOU	1788	O	LEU	A	250	3975	3852	3929	501	794	831	O
ATOM	1789	CB	LEU	A	250	47.973	12.854	8.495	1.00	34.85	C	
ANISOU	1789	CB	LEU	A	250	4409	4465	4358	527	521	744	C
ATOM	1790	CG	LEU	A	250	47.818	13.919	9.554	1.00	40.16	C	
ANISOU	1790	CG	LEU	A	250	5074	5074	5110	530	532	751	C
ATOM	1791	CD1	LEU	A	200	47.274	15.286	9.045	1.00	37.55	C	
ANISOU	1791	CD1	LEU	A	250	4783	4723	4754	577	555	836	C
ATOM	1792	CD2	LEU	A	250	46.977	13.415	10.771	1.00	34.54	C	
ANISOU	1792	CD2	LEU	A	250	4375	4425	4475	523	574	719	C
ATOM	1793	N	VAL	A	251	51.093	12.933	8.162	1.00	31.64	N	
ANISOU	1793	N	VAL	A	251	4302	3965	4034	455	750	716	N
ATOM	1794	CA	VAL	A	251	52.354	18.417	8.732	1.00	34.03	C	
ANISOU	1794	CA	VAL	A	251	4288	4225	4415	416	801	703	C
ATOM	1795	C	VAL	A	251	53.289	13.808	7.616	1.00	36.76	C	
ANISOU	1795	C	VAL	A	251	4654	4575	4737	409	875	735	C
ATOM	1796	O	VAL	A	251	53.937	14.310	7.722	1.00	36.49	O	
ANISOU	1796	O	VAL	A	251	4525	4487	4750	385	901	769	O
ATOM	1797	CB	VAL	A	251	52.985	12.385	9.724	1.00	34.00	C	
ANISOU	1797	CB	VAL	A	251	4277	4251	4505	381	770	525	C
ATOM	1798	CG1	VAL	A	251	54.302	12.935	10.330	1.00	32.24	C	
ANISOU	1798	CG1	VAL	A	251	3992	3929	4328	342	818	509	C
ATOM	1799	CG2	VAL	A	251	51.911	12.140	10.792	1.00	31.69	C	
ANISOU	1799	CG2	VAL	A	251	3934	3913	4183	388	702	605	C
ATOM	1800	N	ASN	A	252	53.255	13.036	6.539	1.00	34.58	N	
ANISOU	1800	N	ASN	A	252	4443	4415	4433	426	878	733	N
ATOM	1801	CA	ASN	A	252	54.052	13.562	5.372	1.00	40.62	C	
ANISOU	1801	CA	ANN	A	252	5182	5140	5110	422	955	774	C
ATOM	1802	C	ASN	A	252	53.639	14.342	4.800	1.00	37.40	C	
ANISOU	1802	C	ASN	A	252	4826	4737	4677	442	922	367	C
ATOM	1803	O	ASN	A	252	54.513	15.692	4.338	1.00	35.95	O	
ANISOU	1803	O	ASN	A	252	4733	4623	4632	419	1053	908	O
ATOM	1804	CB	ADN	A	252	54.097	12.515	4.258	1.00	37.67	C	
ANISOU	1804	CB	ASN	A	252	4617	4646	4649	439	951	745	C
ATOM	1805	CG	ASN	A	252	54.325	11.234	4.725	1.00	35.45	C	
ANISOU	1805	CG	ANN	A	252	4515	4604	4428	417	331	657	C
ATOM	1806	OD1	ASN	A	252	55.485	11.230	5.773	1.00	34.15	O	

TABLE 3-continued

ANISOU	1806	OD1	ASN	A	252	4287	4356	4328	386	931	521	O
ATOM	1807	ND2	ASN	A	252	54.649	10.105	3.937	1.00	36.34	N	
ANISOU	1807	ND2	ASN	A	252	4608	4752	4445	437	905	518	N
ATOM	1808	N	SER	A	253	52.333	15.236	4.756	1.00	41.31	N	
ANISOU	1808	N	SER	A	253	5348	5214	5133	485	941	203	N
ATOM	1809	CA	SER	A	253	51.872	16.595	4.404	1.00	41.99	C	
ANISOU	1809	CA	SER	A	253	5484	5262	5209	511	966	990	C
ATOM	1810	C	SER	A	253	52.376	17.605	5.352	1.00	41.20	C	
ANISOU	1810	C	SER	A	253	5376	5070	5208	462	997	1003	C
ATOM	1811	O	SER	A	253	52.865	18.671	4.907	1.00	41.39	O	
ANISOU	1811	O	SER	A	253	5437	5347	5242	473	1059	1066	O
ATOM	1812	CB	SER	A	253	50.373	16.727	4.310	1.00	38.43	C	
ANISOU	1812	CB	SER	A	253	5053	4840	4709	558	899	1018	C
ATOM	1813	OG	SER	A	253	49.953	15.683	3.525	1.00	38.78	O	
ANISOU	1813	OG	SER	A	253	5596	4969	4568	587	863	994	O
ATOM	1814	N	LEU	A	254	52.290	17.304	6.545	1.00	39.93	N	
ANISOU	1814	N	LEU	A	254	5173	4851	5119	465	956	943	N
ATOM	1815	CA	LEU	A	254	52.337	18.213	7656	1.00	39.22	C	
ANISOU	1815	CA	LEU	A	254	5072	4702	5327	434	981	942	C
ATOM	1816	C	LEU	A	254	54.325	18.525	7.353	1.00	43.12	C	
ANISOU	1816	C	LEU	A	254	5179	4783	5281	353	3051	947	C
ATOM	1817	O	LEU	A	254	54.752	19.704	7.391	1.00	138.20	O	
ANISOU	1817	O	LEU	A	254	4961	4459	5054	350	1130	995	O
ATOM	1818	CB	LEU	A	254	52.708	17.547	9.373	1.00	35.19	C	
ANISOU	1818	CB	LEU	A	254	4533	4178	4583	418	928	868	C
ATOM	1819	CG	LEU	A	254	51.411	17.875	9.827	1.00	38.60	C	
ANISOU	1819	CG	LEU	A	254	4947	4505	5115	457	858	857	C
ATOM	1820	CD1	LEU	A	254	51.203	16.744	10.841	1.00	43.41	C	
ANISOU	1820	CD1	LEU	A	254	5507	5244	5742	442	810	790	C
ATOM	1821	CD2	LEU	A	254	51.327	10.312	13.391	1.00	37.51	C	
ANISOU	1821	CD2	LEU	A	254	4834	4383	5037	464	890	904	C
ATOM	1822	N	LEU	A	255	55.105	17.473	1.095	1.00	36.01	N	
ANISOU	1822	N	LEU	A	255	4622	4312	4747	355	1073	595	N
ATOM	1823	CA	LEU	A	255	56.519	37.524	6.839	1.00	35.55	C	
ANISOU	1823	CA	LEU	A	255	4544	4238	4729	335	1146	889	C
ATOM	1824	C	LEU	A	255	55.725	15.442	5.558	1.00	43.18	C	
ANISOU	1824	C	LEU	A	255	5183	4827	5253	357	1218	974	C
ATOM	1825	O	LEU	A	255	57.534	19.249	5.452	1.00	36.45	O	
ANISOU	1825	O	LEU	A	255	4773	4370	4896	253	1267	1005	O
ATOM	1826	CB	LEU	A	255	57.178	16.240	6.724	1.00	35.65	C	

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ANISOU	1826	CB	LEU	A	255	4634	4439	4855	203	1337	814	C
ATOM	1827	CG	LEU	A	255	57.507	15.577	5.131	1.00	35.79	C	
ANISOU	1827	CG	LEU	A	255	4469	4300	4829	269	1089	736	C
ATOM	1828	CD1	LEU	A	255	57.667	14.174	8.091	1.00	37.35	C	
ANISOU	1828	CD1	LEU	A	255	4633	4558	5000	278	1049	664	C
ATOM	1829	CD2	LEU	A	255	58.808	16.329	8.675	1.00	37.17	C	
ANISOU	1829	CD2	LEU	A	255	4608	4415	5090	213	1142	723	C
ATOM	1830	N	GLU	A	256	55.833	18.295	4.619	1.00	40.57	N	
ANISOU	1830	N	GLU	A	256	5275	4929	5203	356	1198	1015	N
ATOM	1831	CA	GLU	A	256	55.949	19.378	3.387	1.00	42.99	C	
ANISOU	1831	CA	GLU	A	256	5633	5241	5453	351	1254	1104	C
ATOM	1832	C	GLU	A	256	55.550	20.532	3.531	1.00	44.07	C	
ANISOU	1832	C	GLU	A	256	5824	5290	5529	347	1275	1380	C
ATOM	1833	O	GLU	A	256	56.178	21.415	3.094	1.00	43.97	O	
ANISOU	1833	O	GLU	A	256	5345	5239	5623	335	1347	1243	O
ATOM	1834	CB	GLU	A	256	55.108	18.490	2.322	1.00	45.61	C	
ANISOU	1834	CB	GLU	A	256	6006	5657	5565	414	1234	1123	C
ATOM	1835	CG	GLU	A	256	55.153	19.281	1.044	1.00	53.52	C	
ANISOU	1835	CG	GLU	A	256	7088	6532	5602	425	1294	1218	C
ATOM	1836	CD	GLU	A	256	55.319	18.349	-0.097	1.00	72.50	C	
ANISOU	1836	CD	GLU	A	256	9485	9173	8588	433	1310	1202	C
ATOM	1837	OE1	GLU	A	256	56.392	17.703	-0.162	1.00	82.05	O	
ANISOU	1837	OE1	GLU	A	256	10655	10409	10115	400	1355	1148	O
ATOM	1838	OE2	GLU	A	256	54.377	18.250	-0.912	1.00	81.00	O	
ANISOU	1838	OE2	GLU	A	256	10510	15301	9656	491	1274	1240	O
ATOM	1839	N	SER	A	257	54.529	20.809	4.457	1.00	42.14	N	
ANISOU	1839	N	SER	A	257	5585	5012	5414	403	1207	1174	N
ATOM	1840	CA	SER	A	257	54.105	22.205	4.552	1.00	45.40	C	
ANISOU	1840	CA	SER	A	257	6049	5338	5862	413	1215	1244	C
ATOM	1841	C	SER	A	257	55.027	22.914	5.617	1.00	48.06	C	
ANISOU	1841	C	SER	A	257	6386	5583	6318	359	1251	1225	C
ATOM	1842	O	SER	A	257	55.204	24.122	5.521	1.00	45.53	O	
ANISOU	1842	O	SER	A	257	6039	5180	6030	346	1290	1288	O
ATOM	1843	CB	SER	A	257	52.655	22.332	5.112	1.00	48.56	C	
ANISOU	1843	CB	SER	A	257	6463	5733	6248	486	1135	1247	C
ATOM	1844	OG	SER	A	257	51.815	21.518	4.339	1.00	45.74	O	
ANISOU	1844	OG	SER	A	257	6112	5476	5791	534	1091	1243	O
ATOM	1845	N	TYR	A	258	55.615	22.170	6.554	1.00	45.34	N	
ANISOU	1845	N	TYR	A	258	5952	5244	6039	322	1234	1137	N
ATOM	1846	CA	TYR	A	258	56.330	22.753	7.654	1.00	42.66	C	

TABLE 3-continued

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ANISOU	1846	CA	TYR	A	258	5584	4819	5804	279	1247	1103	C
ATOM	1847	C	TYR	A	258	57.645	22.004	7.785	1.00	42.78	C	
ANISOU	1847	C	TYR	A	258	5535	4863	5855	212	1253	1043	C
ATOM	1848	O	TYR	A	258	57.551	21	257	8.783	1.00	41.37	O
ANISOU	1848	O	TYR	A	258	5300	4595	5725	199	1240	950	O
ATOM	1849	CB	TYR	A	258	55.475	22.689	8.946	1.00	42.57	C	
ANISOU	1849	CB	TYR	A	258	5556	4734	5835	303	1171	1053	C
ATOM	1850	CG	TYR	A	258	54.203	23.534	8.830	1.00	46.92	C	
ANISOU	1850	CG	TYR	A	258	6164	5207	6350	367	1139	1110	C
ATOM	1851	CD1	TYR	A	258	54.259	24.910	8.894	1.00	44.73	C	
ANISOU	1851	CD1	TYR	A	258	5042	4936	6195	363	1171	1157	C
ATOM	1852	CD2	TYR	A	258	52.935	22.904	5.609	1.00	44.03	C	
ANISOU	1852	CD2	TYR	A	258	5802	5010	5917	433	1073	1110	C
ATOM	1853	CE1	TYR	A	258	53.131	25.679	8.762	1.00	51.11	C	
ANISOU	1853	CE1	TYR	A	258	6797	5708	6913	429	1141	1220	C
ATOM	1854	CE2	TYR	A	258	51.772	23.797	8.477	1.00	44.73	C	
ANISOU	1854	CE2	TYR	A	258	5933	5075	5983	497	1049	1163	C
ATOM	1855	CZ	TYR	A	258	51.909	25.052	6.554	1.00	48.05	C	
ANISOU	1855	CZ	TYR	A	258	6407	5895	6453	497	1081	1217	C
ATOM	1855	OH	TYR	A	258	50.846	25.926	8.402	1.00	52.22	O	
ANISOU	1856	OH	TYR	A	258	5985	5892	5965	585	1054	1271	O
ATOM	1857	N	PRO	A	259	58.515	22.198	6.779	1.00	45.21	N	
ANISOU	1857	N	PRO	A	259	5351	5186	6140	178	1361	1334	N
ATOM	1858	CA	PRO	A	259	59.839	21.554	5.757	1.00	43.79	C	
ANISOU	1858	CA	PRO	A	259	5607	5038	5995	122	1405	1030	C
ATOM	1859	C	PRO	A	259	60.652	21.688	8.057	1.00	41.90	C	
ANISOU	1859	C	PRO	A	259	5308	4739	5873	71	1395	965	C
ATOM	1860	O	PRO	A	259	61.537	20.855	8.280	1.00	40.85	O	
ANISOU	1860	O	PRO	A	259	5108	4645	5767	42	1401	698	O
ATOM	1861	CB	PRO	A	259	60.572	22.171	5.533	1.00	47.54	C	
ANISOU	1861	CB	PRO	A	259	6110	5513	5436	58	1504	1105	C
ATOM	1862	CG	PRO	A	259	59.619	23.133	4.911	1.00	51.07	C	
ANISOU	1862	CG	PRO	A	259	6645	5530	5829	126	1506	1202	C
ATOM	1863	OD	PRO	A	259	58.267	22.989	5.545	1.00	47.90	C	
ANISOU	1863	CD	PRO	A	259	6264	5523	6413	193	1413	1186	C
ATOM	1864	N	GLU	A	260	60.245	22.655	5.928	1.00	43.76	N	
ANISOU	1864	N	GLU	A	260	5565	4885	4175	63	1373	975	N
ATOM	1865	CA	GLU	A	260	61.017	22.715	10.237	1.00	45.59	C	
ANISOU	1865	CA	GLU	A	260	5743	5057	8512	19	1350	905	C
ATOM	1866	C	GLU	A	260	60.790	21.423	11.108	1.00	44.94	C	

TABLE 3-continued

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ANISOU	1866	C	GLU	A	260	5509	5039	6429	44	1272	314	C	
ATOM	1867	O	GLU	A	260	61.601	21.125	12.326	1.00	42.78	O		
ANISOU	1867	O	GLU	A	260	5276	4749	6228	5	1258	745	O	
ATOM	1868	CB	GLU	A	260	80.520	24.006	11.007	1.00	51.97	C		
ANISOU	1868	CB	GLU	A	260	6593	5765	7384	14	1336	929	C	
ATOM	1869	CG	GLU	A	260	59.746	23.864	12.272	1.000	51.98	C		
ANISOU	1869	CG	GLU	A	260	7353	7015	3675	52	1251	376	C	
ATOM	1870	CD	GLU	A	260	58.196	24.138	12.167	1.00	77.51	C		
ANISOU	1870	CD	GLU	A	260	9886	6933	10582	129	1203	917	C	
ATOM	1871	OE1	GLU	A	260	57.699	25.045	11.421	1.00	76.71	O		
ANISOU	1871	OE1	GLU	A	260	9849	8846	10452	154	1234	998	O	
ATOM	1872	OE2	GLU	A	260	57.444	23.462	12.923	1.00	73.30	O		
ANISOU	1872	OE2	GLU	A	260	9333	8484	10032	167	1139	866	O	
ATOM	1873	N	TYR	A	261	59.696	20.592	13.847	1.00	37.51	N		
ANISOU	1873	N	TYR	A	261	4689	4153	5406	105	1223	315	N	
ATOM	1874	CA	TYR	A	261	59.433	19.399	11.548	1.00	35.43	C		
ANISOU	1874	CA	TYR	A	261	4508	4071	5231	123	1154	737	C	
ATOM	1875	C	TYR	A	261	60.573	18.379	11.306	1.00	35.34	C		
ANISOU	1875	C	TYR	A	261	4312	3986	5131	97	1172	684	C	
ATOM	1876	O	TYR	A	261	60.627	17.531	12.131	1.00	34.64	O		
ANISOU	1876	O	TYR	A	261	4173	39	1	5071	92	1123	514	O
ATOM	1877	CB	TYR	A	261	58.141	15.730	11.175	1.00	36.44	C		
ANISOU	1877	CB	TYR	A	261	4542	4130	5174	186	1102	751	C	
ATOM	1878	CG	TYR	A	261	55.951	19.533	11.776	1.00	35.73	C		
ANISOU	1878	CG	TYR	A	261	4498	4004	5093	222	1061	778	C	
ATOM	1879	CD1	TYR	A	261	56.873	19.786	13.159	1.00	34.45	C		
ANISOU	1879	CD1	TYR	A	261	4311	3785	4993	210	1022	733	C	
ATOM	1880	CD2	TYR	A	261	55	892	19.885	10.989	1.00	33.64	C	
ANISOU	1880	CD2	TYR	A	261	4913	4386	5385	272	3357	840	C	
ATOM	1881	CE1	TYR	A	261	55.772	20.431	13.711	1.00	35.36	C		
ANISOU	1881	CE1	TYR	A	261	4453	3869	5106	248	987	750	C	
ATOM	1882	CE2	TYR	A	261	54.770	20.555	11.513	1.00	38.03	C		
ANISOU	1882	CE2	TYR	A	261	4364	4275	5310	313	1019	851	C	
ATOM	1883	CZ	TYR	A	261	54.728	23.821	12.863	1.00	38.70	C		
ANISOU	1883	CZ	TYR	A	261	4920	4310	5465	300	937	515	C	
ATOM	1884	OH	TYR	A	261	53.584	21.461	13.335	1.00	39.14	O		
ANISOU	1884	OH	TYR	A	261	5013	4341	5518	347	953	832	O	
ATOM	1885	N	LYS	A	262	61.274	18.098	10.193	1.00	38.53	N		
ANISOU	1885	N	LYS	A	262	4723	4428	5522	78	1245	718	N	
ATOM	1886	CA	LYS	A	262	62.436	17.677	9.941	1.00	40.73	C		

TABLE 3-continued

ANISOU	1886	CA	LYS	A	262	4932	4743	5803	52	1274	656	C
ATOM	1887	C	LYS	A	262	60.537	17.301	10.993	1.00	39.35	O	
ANISOU	1887	C	LYS	A	262	4692	4522	5736	0	1272	608	C
ATOM	1888	O	LYS	A	262	64.254	16.844	11.187	1.00	35.30	O	
ANISOU	1888	O	LYS	A	262	4083	4011	5205	-4	1257	542	O
ATOM	1889	CB	LYS	A	262	63.023	17.979	8.594	1.00	44.53	C	
ANISOU	1889	CB	LYS	A	262	5424	5255	5242	30	1365	717	C
ATOM	1890	CG	LYS	A	262	62.243	17.469	7.405	1.00	44.15	C	
ANISOU	1890	CG	LYS	A	262	5423	5277	6075	87	1357	753	C
ATOM	1891	CD	LYS	A	262	53.033	17.940	6.177	1.00	52.16	C	
ANISOU	1891	CD	LYS	A	262	5571	5443	7183	59	1470	305	C
ATOM	1892	CE	LYS	A	262	52.435	17.331	4.845	1.00	57.87	C	
ANISOU	1892	CE	LYS	A	202	7213	7115	7655	104	1487	845	C
ATOM	1893	NZ	LYS	A	262	63.402	17.683	3.767	1.00	63.92	N	
ANISOU	1893	NZ	LYS	A	262	7970	7922	8394	70	1591	370	N
ATOM	1894	N	ASP	A	253	63.635	18.953	11.555	1.00	37.44	N	
ANISOU	1894	N	ASP	A	263	4459	4199	5535	-35	1282	527	N
ATOM	1895	CA	ASP	A	263	64.614	19.003	12.798	1.00	40.79	C	
ANISOU	1895	CA	ASP	A	263	4135	4585	6002	-81	1252	568	C
ATOM	1896	C	ASP	A	263	64.191	18.082	13.915	1.00	35.51	C	
ANISOU	1896	C	ASP	A	263	4135	3929	5430	-52	1163	494	C
ATOM	1897	O	ASP	A	263	65.104	17.581	14.323	1.00	35.87	O	
ANISOU	1897	O	ASP	A	263	4116	3977	5536	-77	1145	427	O
ATOM	1898	CB	ASP	A	263	64.775	20.400	19.421	1.00	41.55	C	
ANISOU	1898	CB	ASP	A	263	4937	4586	6264	-125	1278	488	C
ATOM	1899	CG	ASP	A	263	65.132	21.460	12.362	1.00	50.17	C	
ANISOU	1899	CG	ASP	A	263	6060	5649	7353	-161	1371	669	C
ATOM	1900	OD1	ASP	A	263	65.644	21.092	11.310	1.00	50.37	O	
ANISOU	1900	OD1	ASP	A	263	6068	5732	7337	-168	1431	689	O
ATOM	1901	OD2	ASP	A	263	64.836	22.634	12.574	1.00	52.98	O	
ANISOU	1901	OD2	ASP	A	263	6457	5920	7735	-178	1384	713	O
ATOM	1902	N	SER	A	264	62.859	17.959	14.208	1.00	37.21	N	
ANISOU	1902	N	SER	A	264	4401	4144	5393	-4	1112	509	N
ATOM	1903	CA	SER	A	264	62.473	16.993	15.240	1.00	34.48	C	
ANISOU	1903	CA	SER	A	264	4037	3816	5248	19	1028	447	C
ATOM	1904	C	SER	A	264	62.707	15.594	14.776	1.00	33.07	C	
ANISOU	1904	C	SER	A	264	3828	3710	5026	41	1010	409	C
ATOM	1905	O	SER	A	264	63.055	14.791	15.591	1.00	32.65	O	
ANISOU	1905	O	SER	A	264	3764	3690	5028	39	957	348	O
ATOM	1906	CB	SER	A	264	60.993	17.077	15.768	1.00	36.28	C	

TABLE 3-continued

ANISOU	1906	CB	SER	A	264	4316	4037	5433	61	970	464	C
ATOM	1907	OG	SER	A	264	60.846	18.369	16.414	1.00	36.41	O	
ANISOU	1907	OG	SER	A	264	4357	3977	5501	44	979	484	O
ATOM	1908	N	VAL	A	265	62.481	15.314	13.499	1.00	32.02	N	
ANISOU	1908	N	VAL	A	265	3717	3628	4823	64	1049	445	N
ATOM	1909	CA	VAL	A	265	62.696	13.920	12.995	1.00	34.47	C	
ANISOU	1909	CA	VAL	A	265	4002	4006	5088	89	1030	403	C
ATOM	1910	C	VAL	A	265	64.153	13.565	13.248	1.00	33.38	C	
ANISOU	1910	C	VAL	A	265	3792	3870	5020	57	1050	346	C
ATOM	1911	O	VAL	A	265	64.473	12.464	13.686	1.00	31.10	O	
ANISOU	1911	O	VAL	A	265	3469	3604	4742	71	1000	32.29	O
ATOM	1912	CB	VAL	A	265	62.372	13.779	11.498	1.00	32.29	C	
ANISOU	1912	CB	VAL	A	265	3760	3785	4725	115	1079	448	C
ATOM	1913	CG1	VAL	A	265	62.976	12.454	10.909	1.00	33.70	C	
ANISOU	1913	CG1	VAL	A	265	3902	4028	4873	133	1078	393	C
ATOM	1914	CG2	VAL	A	265	60.843	13.997	11.207	1.00	33.58	C	
ANISOU	1914	CG2	VAL	A	265	3989	3959	4811	156	1045	497	C
ATOM	1915	N	GLN	A	266	65.055	14.524	13.005	1.00	31.20	N	
ANISOU	1915	N	GLN	A	266	3492	3567	4796	13	1121	367	N
ATOM	1916	CA	GLN	A	266	66.472	14.242	13.154	1.00	31.47	C	
ANISOU	1916	CA	GLN	A	266	3447	3611	4898	-19	1146	314	C
ATOM	1917	C	GLN	A	266	66.815	14.083	14.636	1.00	33.70	C	
ANISOU	1917	C	GLN	A	266	3692	3653	5259	-35	1074	254	C
ATOM	1918	O	GLN	A	266	67.486	13	143	15.024	1.00	32.72	O
ANISOU	1918	O	GLN	A	266	3516	3753	5164	-27	1038	190	O
ATOM	1919	CB	GLN	A	266	67.340	15.371	12.525	1.00	34.85	C	
ANISOU	1919	CB	GLN	A	266	3856	4020	5364	-72	1244	356	C
ATOM	1920	CG	GLN	A	266	68.834	15.137	12.794	1.00	36.53	C	
ANISOU	1920	CG	GLN	A	266	3975	4245	5659	-110	1267	296	C
ATOM	1921	CD	GLN	A	266	69.718	16.214	12.161	1.00	40.23	C	
ANISOU	1921	CD	GLN	A	266	4417	4699	6168	-172	1369	337	C
ATOM	1922	OE1	GLN	A	266	69.404	16.740	11.111	1.00	39.64	O	
ANISOU	1922	OE1	GLN	A	266	4389	4637	6036	-175	1436	407	O
ATOM	1923	NE2	GLN	A	266	70.839	16.491	12.783	1.00	39.04	N	
ANISOU	1923	NE2	GLN	A	266	4191	4528	6115	-222	1378	295	N
ATOM	1924	N	GLU	A	267	66.387	15.042	15.465	1.00	34.02	N	
ANISOU	1924	N	GLU	A	267	3761	3830	5335	-57	1055	275	N
ATOM	1925	CA	GLU	A	267	66.729	14.984	16.858	1.00	34.53	C	
ANISOU	1925	CA	GLU	A	267	3794	3857	5468	-74	990	220	C
ATOM	1926	C	GLU	A	267	66.139	13.749	17.601	1.00	33.60	C	

TABLE 3-continued

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ANISOU 1926 C GLU A 267	3685 3764 5319 -32 897 175 C
ATOM 1927 O GLU A 267	66.832 13.167 18.443 1.00 31.77 O
ANISOU 1927 O GLU A 267	3406 3531 5135 -38 848 114 O
ATOM 1928 CB GLU A 267	66.271 16.253 17.573 1.00 31.70 C
ANISOU 1928 CB GLU A 267	4234 438T 5935 -100 936 248 C
ATOM 1929 CG GLU A 267	66.593 16.110 190.084 1.00 31.41 C
ANISOU 1929 CG GLU A 267	4153 4315 5930 -116 913 156 C
ATOM 1930 CD GLU A 267	66.789 17.543 19.686 1.00 48.73 C
ANISOU 1930 CD GLU A 267	5612 5475 1430 -152 930 195 C
ATOM 1931 OE1 GLU A 267	66.074 18.495 19.269 1.00 45.70 O
ANISOU 1931 OE1 GLU A 267	5285 3055 7024 -160 968 256 O
ATOM 1932 OE2 GLU A 267	67.653 11 653 20.565 1.00 59.54 O
ANISOU 1932 OE2 GLU A 267	6932 6320 8372 -197 902 143 O
ATOM 1933 N THR A 268	64.877 13.405 17.324 1.00 27.89 N
ANISOU 1933 N THR A 268	3021 3050 4517 8 873 206 N
ATOM 1934 CA THR A 268	64.267 32.374 17.372 1.00 28.02 C
ANISOU 1934 CA THR A 268	3049 3102 4496 43 792 171 C
ATOM 1935 C THR A 268	55.055 17.557 17.438 1.00 29.90 C
ANISOU 1935 C THR A 268	3243 3387 4731 61 781 122 C
ATOM 1936 O THR A 268	55.175 9.967 18.243 1.00 32.56 O
ANISOU 1936 O THR A 268	3555 3724 5081 73 710 74 O
ATOM 1937 CB THR A 268	62.762 11.999 17.563 1.00 25.21 C
ANISOU 1937 CB THR A 268	2755 2755 4056 76 771 213 C
ATOM 1938 OG1 THR A 268	62.514 12.066 16.138 1.00 28.75 O
ANISOU 1938 OG1 THR A 268	3230 3255 5550 96 323 257 O
ATOM 1939 CG2 THR A 268	51.019 13.162 18.263 1.00 27.50 C
ANISOU 1939 CG2 THR A 268	3124 3046 4393 70 765 249 C
ATOM 1940 N ALA A 269	65.525 10.579 15.247 1.00 31.39 N
ANISOU 1940 N ALA A 269	3412 3530 4906 53 849 133 N
ATOM 1941 CA ALA A 269	56.475 9.787 15.733 1.00 54.59 C
ANISOU 1941 CA ALA A 269	3783 4072 3224 83 852 83 C
ATOM 1942 C ALA A 269	57.607 9.752 16.455 1.00 33.59 C
ANISOU 1942 C ALA A 269	3581 3930 5200 54 842 27 C
ATOM 1943 O ALA A 269	68.361 8.679 16.734 1.00 32.21 O
ANISOU 1943 O ALA A 269	3344 3747 5135 82 793 -32 O
ATOM 1944 CB ALA A 269	56.742 9.994 14.212 1.00 34.24 C
ANISOU 1944 CB ALA A 269	3726 4060 5223 87 942 114 C
ATOM 1945 N GLU A 270	68.384 10.928 15.579 1.00 31.55 N
ANISOU 1945 N GLU A 270	3321 3563 5316 13 888 44 N
ATOM 1946 CA GLU A 270	69.538 11.031 17.407 1.00 32.33 C

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TABLE 3-continued

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ANISOU	1946	CA	GLU	A	270	3301	3705	5255	-19	377	-9	C
ATOM	1947	C	GLU	A	270	59.595	10.601	18.846	1.00	31.81	C	
ANISOU	1947	C	GLU	A	270	3235	3612	5240	-14	773	-54	C
ATOM	1948	O	GLU	A	270	70.537	9.889	19.322	1.00	29.50	O	
ANISOU	1948	O	GLU	A	270	2981	3332	4997	-5	735	-115	O
ATOM	1949	CB	GLU	A	270	70.223	12.499	17.327	1.00	33.91	C	
ANISOU	1949	CB	GLU	A	270	3229	3623	5276	-81	949	22	C
ATOM	1950	CG	GLU	A	270	70.813	12.770	35.919	1.00	37.26	C	
ANISOU	1950	CG	GLU	A	270	3880	4340	5939	-95	1055	51	C
ATOM	1951	CD	GLU	A	270	71.148	14.247	15.7.50	1.00	45.34	C	
ANISOU	1951	CD	GLU	A	270	4900	5321	7007	-159	1193	99	C
ATOM	1952	OE1	GLU	A	270	70.7139	14.995	16.640	1.00	46.57	O	
ANISOU	1952	OE1	GLU	A	270	5852	5581	7460	-181	1093	310	O
ATOM	1953	OE2	GLU	A	270	71.684	14	587	14.718	1.00	45.33	O
ANISOU	1953	OE2	GLU	A	270	5305	5470	7128	-185	1220	1131	O
ATOM	1954	N	VAL	A	271	68.516	10.892	19.552	1.00	29	03	N
ANISOU	1954	N	VAL	A	271	2945	3226	4858	-12	783	-28	N
ATOM	1955	CA	VAL	A	271	88.584	10.854	20.91	1.00	30.47	C	
ANISOU	1955	CA	VAL	A	271	3122	3379	5077	-16	647	-73	C
ATOM	1956	C	VAL	A	271	58.356	9.163	21.370	1.00	29.52	C	
ANISOU	1956	C	VAL	A	271	3011	3282	4924	29	566	-106	C
ATOM	1957	O	VAL	A	271	69.557	8.780	22.575	1.00	32.17	O	
ANISOU	1957	O	VAL	A	271	3338	3597	5287	30	487	-144	O
ATOM	1958	CB	VAL	A	271	67.543	11.582	21	824	1.00	28.72	C
ANISOU	1958	CB	VAL	A	271	2957	3111	4845	-34	525	-39	C
ATOM	1959	CG1	VAL	A	271	67.9799	13.093	21.634	1.00	30.67	C	
ANISOU	1959	CG1	VAL	A	271	3195	3318	5141	-81	695	-10	C
ATOM	1960	CG2	VAL	A	271	86.146	11.244	21.574	1.00	28.94	C	
ANISOU	1960	CG2	VAL	A	271	3059	3152	4784	1	610	4	C
ATOM	1961	N	VAL	A	272	67.912	8.357	20.407	1.00	29.72	N	
ANISOU	1961	N	VAL	A	272	3060	3345	4839	65	580	-94	N
ATOM	1962	CA	VAL	A	272	67.761	6.915	20.658	1.00	31.35	C	
ANISOU	1962	CA	VAL	A	272	3276	3567	5370	105	506	-130	C
ATOM	1963	C	VAL	A	272	69.527	5.105	23.253	1.00	31.76	C	
ANISOU	1964	C	VAL	A	272	3260	3545	5163	128	508	-186	C
ATOM	1964	O	VAL	A	272	69.132	4.891	20.530	1.00	32.93	O	
ANISOU	1964	O	VAL	A	272	3410	3797	5305	163	442	-224	O
ATOM	1965	CB	VAL	A	272	66.500	5.321	20.048	1.00	30.43	C	
ANISOU	1965	CB	VAL	A	272	3277	3521	4915	133	498	-95	C
ATOM	1966	CG1	VAL	A	272	65.277	6.963	20.752	1.00	28.83	C	

TABLE 3-continued

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ANISOU	1966	CG1	VAL	A	272	3082	3243	4529	116	478	-52	C
ATOM	1967	CG2	VAL	A	272	65.544	6.459	18.501	1.00	28.45	C	
ANISOU	1967	CG2	VAL	A	272	2974	3250	4576	144	561	-72	C
ATOM	1968	N	ILE	A	273	70.905	6.770	19.638	1.00	31.66	N	
ANISOU	1968	N	ILE	A	273	3186	3643	5195	157	583	-192	N
ATOM	1969	CA	ILE	A	273	71.301	6.108	19.291	1.00	30.55	C	
ANISOU	1969	CA	ILE	A	273	2966	3538	5104	128	532	-251	C
ATOM	1970	C	ILE	A	273	71.929	5.427	20.498	1.00	32.51	C	
ANISOU	1970	C	ILE	A	273	3192	3771	5421	142	491	-308	C
ATOM	1971	O	ILE	A	273	72.372	4.229	20.398	1.00	36.70	O	
ANISOU	1971	O	ILE	A	273	3588	4312	5943	193	454	-356	O
ATOM	1972	CB	ILE	A	273	72.302	7.054	16.538	1.00	29.95	C	
ANISOU	1972	CB	ILE	A	273	2320	3484	5075	91	693	-248	C
ATOM	1973	CG1	ILL	A	273	71.713	7.458	17.203	1.00	29.99	C	
ANISOU	1973	CG1	ILL	A	273	2565	3518	5011	89	782	-192	C
ATOM	1974	CG2	ILE	A	273	73.711	6.401	18.378	1.00	32.11	C	
ANISOU	1974	CG2	ILE	A	273	2995	3792	5411	112	695	-317	C
ATOM	1975	CD1	ILE	A	273	71.543	6.328	15.149	1.00	30.91	C	
ANISOU	1975	CD1	ILE	A	273	2997	3664	5063	144	793	-210	C
ATOM	1976	N	PRO	A	274	71.903	6.079	21.577	1.00	34.25	N	
ANISOU	1976	N	PRO	A	274	3402	3945	5665	108	454	-308	N
ATOM	1977	CA	PRO	A	274	72.591	5.352	22.785	1.00	36.60	C	
ANISOU	1977	CA	PRO	A	274	3665	4225	8312	127	359	-362	C
ATOM	1978	C	PRO	A	274	71.777	4.155	23.298	1.00	35.89	C	
ANISOU	1978	C	PRO	A	274	3643	4124	5868	168	279	-354	C
ATOM	1979	O	PRO	A	274	72.252	3.441	24.202	1.00	35.7	O	
ANISOU	1979	O	PRO	A	274	3604	4386	5875	190	184	-404	O
ATOM	1980	CB	PRO	A	274	72.649	6.383	23.930	1.00	33.69	C	
ANISOU	1980	CB	PRO	A	274	3927	4455	6318	73	334	-358	C
ATOM	1981	CG	PRO	A	274	72.593	7.703	23.244	1.00	37.55	C	
ANISOU	1981	CG	PRO	A	274	3719	4245	6114	32	432	-314	C
ATOM	1982	CD	PRO	A	274	71.572	7.479	22.068	1.00	36.93	C	
ANISOU	1982	CD	PRO	A	274	3763	4254	6019	54	488	-266	C
ATOM	1983	N	LEU	A	275	70.524	3.387	22.688	1.00	33.93	N	
ANISOU	1983	N	LEU	A	275	3466	3883	5542	180	288	-321	N
ATOM	1984	CA	LEU	A	275	59.792	2.729	23.110	1.00	30.54	C	
ANISOU	1984	CA	LEU	A	275	3116	3452	5073	211	207	-319	C
ATOM	1985	C	LEU	A	275	59.978	1.552	22.319	1.00	32.57	C	
ANISOU	1985	C	LEU	A	275	3357	3721	5296	250	210	-340	C
ATOM	1986	O	LEU	A	275	690.277	0.515	22.225	1.00	31.30	O	

TABLE 3-continued

ANISOU	1986	O	LEU	A	275	3254	3548	5090	285	151	-344	O
ATOM	1987	CB	LEU	A	275	68.277	3.135	23.245	1.00	29.57	C	
ANISOU	1987	CB	LEU	A	275	3003	3307	487	188	212	-258	C
ATOM	1988	CG	LEU	A	275	67.950	4	512	23.856	1.00	29.65	C
ANISOU	1988	CG	LEU	A	275	3.082	3300	4889	143	238	-226	C
ATOM	1989	CD1	LEU	A	275	60.450	4.700	23.90	1.00	30.84	C	
ANISOU	1989	CD1	LEU	A	275	3305	3446	4965	105	239	-173	C
ATOM	1990	CD2	LEU	A	275	68.512	4.424	25.279	1.00	30.94	C	
ANISOU	1990	CD2	LEU	A	275	8225	3433	5097	134	150	-200	C
ATOM	1991	N	MET	A	276	70.	953	1.595	21.210	1.00	31.88	N
ANISOU	1991	N	MET	A	276	3203	3007	5244	273	275	-875	
ATOM	1992	CA	MET	A	276	71.193	0.633	20.196	1.00	35.63	C	
ANISOU	1992	CA	MET	A	276	3678	4176	5704	324	255	-409	C
ATOM	1993	C	MET	A	276	72.293	-0.249	20.725	1.00	36.41	C	
ANISOU	1993	C	MET	A	276	3716	4258	5860	365	220	-475	C
ATOM	1994	O	MET	A	276	73.027	0.186	21.569	1.00	37.20	O	
ANISOU	1994	O	MET	A	276	3765	4346	6023	348	195	-494	O
ATOM	1995	CB	MET	A	276	71.578	1.230	18.804	1.00	35.30	C	
ANISOU	1995	CB	MET	A	276	3555	4152	5020	319	400	-402	C
ATOM	1996	CG	MET	A	276	70.533	2.192	18.244	1.00	37.77	C	
ANISOU	1996	CG	MET	A	276	3956	4498	5595	232	466	-332	C
ATOM	1997	SD	MET	A	276	69.059	1	209	17.906	1.00	33.29	S
ANISOU	1997	SD	MET	A	276	4122	4560	5866	310	417	-308	S
ATOM	1998	CE	MET	A	276	57.855	2.472	18.432	1.00	41.58	C	
ANISOU	1998	CE	MET	A	276	4593	4955	5251	257	430	-230	C
ATOM	1999	N	ALA	A	277	72.443	-1.484	20.216	1.00	34.03	N	
ANISOU	1999	N	ALA	A	277	3423	3902	5545	422	190	-515	N
ATOM	2000	CA	ALA	A	277	73.442	-2.397	20.751	1.00	40.65	C	
ANISOU	2000	CA	ALA	A	277	4241	4811	6470	471	118	-579	C
ATOM	2001	C	ALA	A	277	74.790	-1.891	20.303	1.00	47.10	C	
ANISOU	2001	C	ALA	A	277	4920	5545	7320	475	153	-623	C
ATOM	2002	O	ALA	A	277	75.503	2.111	20.076	1.00	50.59	O	
ANISOU	2002	O	ALA	A	277	5297	6082	7644	497	135	-670	O
ATOM	2003	CB	ALA	A	277	73.256	-3.809	20.238	1.00	41.34	C	
ANISOU	2003	CB	ALA	A	277	4305	4825	6463	533	75	-613	C
ATOM	2004	N	ASN	A	278	74.752	-1.153	19.205	1.00	44.86	N	
ANISOU	2004	N	ASN	A	278	4517	5426	7020	452	293	-602	N
ATOM	2005	CA	ASN	A	278	75.704	-1.328	18.134	1.00	57.59	C	
ANISOU	2005	CA	ASN	A	278	74313	3353	9933	485	307	-649	C
ATOM	2006	C	ASN	A	278	75.507	-2.649	17.340	1.00	69.65	C	

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TABLE 3-continued

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ANISOU	2006	C ASN A	278	7704	8613	10140	653	350	-695	C
ATOM	2007	O ASN A	278	75.217	-3.720	17.702	1.00	74.00	O	
ANISOU	2007	O ASN A	278	3238	9142	10736	517	271	-756	O
ATOM	2008	CB ASN A	278	77.033	-0.852	18.509	1.00	50.20	C	
ANISOU	2008	CB ASN A	278	7006	8053	9713	475	375	-689	C
ATOM	2009	CG ASN A	278	75.995	8053	18.714	1.00	52.21	C	
ANISOU	2009	CG ASN A	278	6607	7630	9349	390	444	-539	C
ATOM	2010	OD1 ASN A	278	76.971	1.285	17.690	1.00	58.53	O	
ANISOU	2010	OD1 ASN A	278	5130	7253	3856	370	549	-612	O
ATOM	2011	ND2 ASN A	278	75.800	1.155	19.915	1.00	65.37	N	
ANISOU	2011	ND2 ASN A	278	7024	8037	9778	357	384	-618	N
ATOM	2012	OXT ASN A	278	75.073	-2.576	10.261	1.00	71.58	O	
ANISOU	2012	OXT ASN A	278	8002	8899	10333	556	417	-572	O
TER	2013	ASN A	278							
ATOM	2014	N ASN B	6	50.962	15.155	4.943	1.00	73.54	N	
ANISOU	2014	N ASN B	6	9137	8979	9824	-681	1694	-358	N
ATOM	2015	CA ASN B	6	82.343	16.416	5.467	1.00	77.214	C	
ANISOU	2015	CA ASN B	6	9559	9.515	10292	-859	1762	-495	C
ATOM	2016	C ASN B	6	82.968	15.793	6.360	1.00	74.00	C	
ANISOU	2016	C ASN B	6	8972	9252	9894	-872	1638	-624	C
ATOM	2017	O ASN B	6	84.199	15.190	6.418	1.00	72.84	O	
ANISOU	2017	O ASN B	6	8731	9226	9719	-986	1660	-728	O
ATOM	2018	CB ASN B	6	82.445	17.841	6.084	1.00	72.41	C	
ANISOU	2018	CB ASN B	6	9079	3732	9703	-989	19429	-499	C
ATOM	2019	CG ASN B	6	83.503	17.951	7.174	1.00	72.65	C	
ANISOU	2019	CG ASN B	6	9033	8831	9741	-1164	1954	-558	C
ATOM	2020	OD1 ASN B	6	83.383	17.335	8.242	1.00	68.47	O	
ANISOU	2020	OD1 ASN B	6	8410	8356	9240	-1142	1853	-722	O
ATOM	2021	ND2 ASN B	6	84.533	18.762	6.924	1.00	73.29	N	
ANISOU	2021	ND2 ASN B	6	9776	9548	13424	-1348	2092	-721	N
ATOM	2022	N LEU B	7	82.130	14.462	7.332	1.50	62.57	N	
ANISOU	2022	N LEU B	7	7431	7805	5489	-754	1514	-611	N
ATOM	2023	CA LEU B	7	82.650	13.233	7.727	1.05	82.74	C	
ANISOU	2023	CA LEU B	7	7349	7968	8021	-727	1383	-708	C
ATOM	2024	C LEU B	7	83.409	12.244	6.755	1.00	63.86	C	
ANISOU	2024	C LEU B	7	7380	8262	8623	-507	1332	-742	C
ATOM	2025	O LEU B	7	84.339	11.521	7.175	1.00	53.00	O	
ANISOU	2025	O LEU B	7	7139	8235	8511	-708	1280	-832	O
ATOM	2026	CB LEU B	7	81.539	12.494	8.480	1.00	57.54	C	
ANISOU	2026	CB LEU B	7	5685	7263	7913	-633	1280	-654	C

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ATOM	2027	CG	LEU	B	7	81.809	11.000	8.002	1.00	57.28	C
ANISOU	2027	CG	LEU	B	7	5525	7354	7886	-513	1143	-711 C
ATOM	2028	CD1	LEU	B	7	82.320	10.741	9.607	1.00	57.57	C
ANISOU	2028	CD1	LEU	B	7	7733	8782	5193	-590	1125	-812 C
ATOM	2029	CD2	LEU	B	7	80.561	13.225	8.965	1.00	56.39	C
ANISOU	2029	CD2	LEU	B	7	6489	7245	7882	-394	1057	-554 C
ATOM	2030	N	LEU	B	8	82.939	12.222	5.503	1.00	60.17	N
ANISOU	2030	N	LEU	B	8	5966	7775	8120	-523	1345	-670 N
ATOM	2031	CA	LEU	B	8	39.575	11.521	4.419	1.00	58.45	C
ANISOU	2031	CA	LEU	B	8	6677	7384	7849	-609	1322	-703 C
ATOM	2032	C	LEU	B	8	85.048	12.153	4.198	1.00	51.00	C
ANISOU	2032	C	LEU	B	8	6957	3385	3134	-755	1443	-781 C
ATOM	2033	O	LEU	B	8	86.056	11.430	4.154	1.00	53.55	O
ANISOU	2033	O	LEU	B	8	7146	3553	3448	-764	1417	-672 O
ATOM	2034	CB	LEU	B	8	82.843	11.4B5	3.127	1.00	53.74	C
ANISOU	2034	CB	LEU	B	8	6151	7356	7200	-528	1327	-513 C
ATOM	2035	CG	LEU	B	8	81.579	10.592	2.121	1.00	61.25	C
ANISOU	2035	CG	LEU	B	8	7114	7931	8132	-392	1201	-556 C
ATOM	2036	CD1	LEU	B	8	83.957	10.285	1.825	1.00	51.99	C
ANISOU	2036	CD1	LEU	B	8	7246	8108	8200	-324	1173	-535 C
ATOM	2037	CD2	LEU	B	8	81.822	9.234	3.949	1.30	62.14	C
ANISOU	2037	CD2	LEU	B	8	7112	8154	8343	-340	1099	-654 C
ATOM	2038	N	VAL	B	9	85.060	13.492	4.377	1.30	56.85	N
ANISOU	2038	N	VAL	B	9	6547	7450	7593	-854	1573	-733 N
ATOM	2039	CA	VAL	B	9	36.273	14.286	3.875	1.00	51.59	C
ANISOU	2039	CA	VAL	B	9	7144	8124	8173	-1333	1705	-803 C
ATOM	2040	C	VAL	B	9	87.224	43.974	5.318	1 00	65.59	C
ANISOU	2040	C	VAL	B	9	7487	8738	8695	-1115	1657	-932 C
ATOM	2041	O	VAL	B	9	88.421	13.735	4.797	1.00	52.67	O
ANISOU	2041	O	VAL	B	9	6987	8525	8299	-1193	1692	-1023 O
ATOM	2042	CB	VAL	B	9	85.307	15.826	3.755	1.00	52.00	C
ANISOU	2042	CB	VAL	B	9	7483	8118	8331	-1146	1863	-733 C
ATOM	2043	CG1	VAL	B	9	37.281	15.589	3.288	1.00	50.15	C
ANISOU	2043	CG1	VAL	B	9	7118	7312	7925	-1339	2019	-799 C
ATOM	2044	CG2	VAL	B	9	34.638	15.089	2.741	1.00	53.57	C
ANISOU	2044	CG2	VAL	B	9	7705	6077	8372	-1024	1377	-577 C
ATOM	2045	N	ARG	B	10	35.632	13.933	6.238	1.00	63.22	N
ANISOU	2045	N	ARG	B	10	7195	8380	3445	-1095	1602	-938 N
ATOM	2046	CA	ARG	B	10	87.473	13.548	7.401	1.00	63.13	C
ANISOU	2046	CA	ARG	B	10	7046	8507	8452	-1152	1540	-1047 C

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TABLE 3-continued

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ATOM	2047	C ARG B 10	87.943	12.083	7.293	1.00	58.70	C
ANISOU	2047	C ARG B 10	6311	3106	7888	-1022	1413	-1384 C
ATOM	2048	O ARG B 10	39.120	11.806	7.530	1.00	53.54	O
ANISOU	2048	O ARG B 10	6304	6931	6521	-1080	1405	-1175 O
ATOM	2049	CB ARG B 10	86.762	13.880	8.724	1.00	65.65	C
ANISOU	2049	CB ARG B 10	7550	8846	8927	-1153	1510	-1041 C
ATOM	2050	CG ARG B 10	67.715	14.1145	9.906	1.00	75.53	C
ANISOU	2050	CG ARG B 10	8699	10213	10163	-1313	1530	-1162 C
ATOM	2051	CD ARG B 10	67.013	14.536	11.217	1.00	76.05	C
ANISOU	2051	CD ARG B 10	8719	10058	11116	-1330	1496	-1166 C
ATOM	2052	NE ARG B 10	86.3615	13.394	11.882	1.00	74.71	N
ANISOU	2052	NE ARG B 10	8500	9914	9973	-1170	1345	-1127 N
ATOM	2053	CZ ARG B 10	85.038	13.208	11.996	1.00	73.25	C
ANISOU	2053	CZ ARG B 10	9056	13210	10455	-1043	1315	-1034 C
ATOM	2054	NH1 ARG B 10	84.587	12.126	12.625	1.00	68.61	N
ANISOU	2054	NH1 ARG B 10	7732	9025	9252	-912	1138	-1010 N
ATOM	2055	NH2 ARG B 10	84.150	14.064	11.483	1.00	81.15	N
ANISOU	2055	NH2 ARG B 10	9662	10466	10935	-1042	1415	-958 N
ATOM	2056	N LEU B 11	67.070	11.153	6.890	1.00	52.29	N
ANISOU	2056	N LEU B 11	5527	7246	7096	-850	1325	-1317 N
ATOM	2057	CA LEU B 11	87.531	9.759	6.701	1.03	52.21	C
ANISOU	2057	CA LEU B 11	5382	7361	7053	-723	1229	-1054 C
ATOM	2058	C LEU B 11	88.733	9.536	5.742	1.00	56.38	C
ANISOU	2058	C LEU B 11	5765	7999	7544	-761	1292	-1120 C
ATOM	2059	O LEU B 11	89.617	8.830	6.035	1.00	59.87	O
ANISOU	2059	O LEU B 11	6023	6555	7953	-722	1248	-1138 O
ATOM	2060	CB LEU B 11	36.304	8.913	6.152	1.00	55.81	C
ANISOU	2060	CB LEU B 11	5913	7724	7573	-565	1155	-982 C
ATOM	2061	CG LEU B 11	66.306	7.379	6.086	1.30	53.17	C
ANISOU	2061	CG LEU B 11	5495	7448	7260	-436	1057	-1006 C
ATOM	2062	CD1 LEU B 11	87.259	6.790	7.156	1.00	55.31	C
ANISOU	2062	CD1 LEU B 11	5750	7964	7583	-393	1301	-1066 C
ATOM	2063	CD2 LEU B 11	84.949	6.331	6.200	1.00	57.55	C
ANISOU	2063	CD2 LEU B 11	6142	7376	7333	-309	980	-936 C
ATOM	2064	N ARG B 12	88.893	10.277	4.605	1.30	55.00	N
ANISOU	2064	N ARG B 12	5715	7820	7361	-327	1398	-1091 N
ATOM	2065	CA ARG B 12	89.741	10.165	3.572	1.00	56.45	C
ANISOU	2065	CA ARG B 12	5316	8133	7499	-366	1477	-1147 C
ATOM	2066	C ARG B 12	91.021	10.712	4.143	1.00	62.35	C
ANISOU	2066	C ARG B 12	6402	9018	8242	-1020	1538	-1240 C

TABLE 3-continued

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ATOM	2067	O	ARG	B	12	92.094	10.035	4.043	1.00	65.52	O
ANISOU	2067	O	ARG	B	12	3670	9513	6553	-1004	1533	-1320 O
ATOM	2068	CB	ARG	B	12	89.412	11.012	2.312	1.00	53.58	C
ANISOU	2068	CB	ARG	B	12	5602	7535	7371	-931	1593	-1331 C
ATOM	2069	CG	ARG	B	12	86.342	10.455	1.407	1.00	53.86	C
ANISOU	2069	CG	ARG	B	12	5741	7645	7073	-796	1542	-1002 C
ATOM	2070	CD	ARG	B	12	68.424	10.661	-0.064	1.00	53.09	C
ANISOU	2070	CD	ARG	B	12	5728	7556	6836	-332	1643	-962 C
ATOM	2071	NE	ARG	B	12	87.530	9.390	-0.749	1.77	57.31	N
ANISOU	2071	NE	ARG	B	12	5302	8076	7396	-685	1550	-927 N
ATOM	2072	CZ	ARG	B	12	87.919	3.699	-1.257	1.00	53.29	C
ANISOU	2072	CZ	ARG	B	12	6465	8430	7532	-603	1511	-1004 C
ATOM	2073	NH1	ARG	B	12	39.240	8.394	-1.247	1.90	94.11	N
ANISOU	2073	NH1	ARG	B	12	55.54	7910	6985	-5134	1564	-1111 N
ATOM	2074	NH2	ARG	B	12	86.980	7.834	-1.797	1.00	54.51	N
ANISOU	2074	NH2	ARG	B	12	5915	7794	7032	-492	1428	-979 N
ATOM	2075	N	SER	B	13	90.881	11.520	4.597	1.00	60.89	N
ANISOU	2075	N	SER	B	13	6335	8746	8055	-1172	1808	-1231 N
ATOM	2076	CA	SER	B	13	91.934	12.670	5.333	1.00	71.19	C
ANISOU	2076	CA	SER	B	13	7544	10157	9349	-1380	1677	-1324 C
ATOM	2077	C	SER	B	13	32.560	11.887	6.499	1.00	78.29	C
ANISOU	2077	C	SER	B	13	8245	11230	10271	-1321	1554	-1403 C
ATOM	2078	O	SER	B	13	93.787	11.333	3.506	1.00	95.60	O
ANISOU	2078	O	SER	B	13	10381	13751	12572	-1404	1575	-1493 O
ATOM	2079	CB	SER	B	13	91.354	13.979	5.843	1.00	39.23	C
ANISOU	2079	CB	SER	B	13	7457	9733	9104	-1504	1751	-1293 C
ATOM	2080	OG	SER	B	13	92.232	15.024	5.584	1.00	75.40	O
ANISOU	2080	OG	SER	B	13	8246	10547	9853	-1723	1915	-1355 O
ATOM	2081	N	ASN	B	14	91.7134	11.260	7.340	1.00	72.08	N
ANISOU	2081	N	ASN	B	14	7527	10415	9557	-1187	1427	-1354 N
ATOM	2082	CA	ASN	B	14	92.223	10.575	3.543	1.00	76.05	C
ANISOU	2082	CA	ASN	B	14	7830	11932	10032	-1139	1807	-1403 C
ATOM	2083	C	ASN	B	14	92.539	9.102	8.330	1.30	77.20	C
ANISOU	2083	C	ASN	B	14	7840	11291	10202	-931	1208	-1398 C
ATOM	2084	O	ASN	B	14	92.002	3.349	9.272	1.03	82.63	O
ANISOU	2084	O	ASN	B	14	8411	12034	10901	-839	3057	-1237 O
ATOM	2085	CB	ASN	B	14	91.222	10.749	9091	1.00	82.14	C
ANISOU	2085	CB	ASN	B	14	3744	11575	13321	-1122	1239	-1350 C
ATOM	2086	CG	ASN	B	14	91.240	12.151	10.271	1.00	84.08	C
ANISOU	2086	CG	ASN	B	14	9046	11858	11041	-1342	1336	-1403 C

TABLE 3-continued

ATOM	2087	OD1	ASN	B	14	93.938	12.339	11.440	1.00	53.17	O
ANISOU	2987	OD1	ASN	B	14	10853	13628	12820	-1070	1237	-1417 O
ATOM	2088	ND2	ASN	B	14	91.603	13.138	5.455	1.09	87.38	N
ANISOU	2088	ND2	ASN	B	14	9525	12235	11441	-1496	1483	-1426 N
ATOM	2089	N	MET	B	15	9252	8.717	7.031	1.00	74.33	N
ANISOU	2089	N	MET	B	15	7503	10903	9843	-860	1258	-1381 N
ATOM	2090	CA	MET	B	15	92.599	7.361	5.709	1.00	75.31	C
ANISOU	2090	CA	MET	B	15	7502	11082	9990	-453	1133	-1370 C
ATOM	2091	C	MET	B	15	94.183	7.013	5.545	1.33	83.33	C
ANISOU	2991	C	MET	B	15	7583	11999	10655	-667	1219	-1465 C
ATOM	2092	O	MET	B	15	94.588	5.326	6.499	1.00	71.59	O
ANISOU	2092	O	MET	B	15	6763	10977	9578	-450	1154	-1474 O
ATOM	2093	CB	MET	B	15	92.142	1.304	5.205	1.00	75.51	C
ANISOU	2093	CB	MET	B	15	7698	10594	9599	-631	1258	-1345 C
ATOM	2094	CG	MET	B	15	91.534	5.014	4.756	1.00	73.83	C
ANISOU	2094	CG	MET	B	15	7522	10706	9812	-434	1200	-1314 C
ATOM	2095	SD	MET	B	15	90.666	6.340	3.182	3.00	65.70	S
ANISOU	2095	SD	MET	B	15	7184	10359	9239	-455	1280	-1270 S
ATOM	2096	CE	MET	B	15	91.829	7.258	2.133	1.00	72.55	C
ANISOU	2096	CE	MET	B	15	7506	10555	9541	-615	1442	-1325 C
ATOM	2097	N	GLU	B	16	94.983	8.077	6.487	1.00	83.34	N
ANISOU	2097	N	GLU	B	16	8252	12445	10958	-871	1312	-1527 N
ATOM	2098	CA	GLU	B	16	90.442	5.001	7.539	1.00	93.59	C
ANISOU	2098	CA	GLU	B	16	9314	14034	12241	-923	1343	-4619 C
ATOM	2099	C	GLU	B	16	97.033	3.379	7.439	1.09	52.96	C
ANISOU	2099	C	GLU	B	16	9027	14103	12139	-748	1208	-1628 C
ATOM	2100	O	GLU	B	16	97.733	5.930	8.940	1.00	89.55	O
ANISOU	2100	O	GLU	B	16	8457	13798	11786	-605	1210	-1651 O
ATOM	2101	CB	GLU	B	16	97.026	9.407	5.825	1.00	39.05	C
ANISOU	2101	CB	GLU	B	16	9981	14767	12885	-1204	1435	-1557 C
TOM	2102	CG	GLU	B	16	98.417	9.661	5.266	1.00104.39	C	
ANISOU	2102	CG	GLU	B	16	10454	15671	12539	-1323	1541	-1784 C
ATOM	2103	CD	GLU	B	16	99.459	8.722	7.853	1.00106.33	C	
ANISOU	2103	CD	GLU	B	16	10410	16192	13300	-1200	1445	-1834 C
ATOM	2104	OE1	GLU	B	16	99.703	3.794	8.079	1.00110.29	O	
ANISOU	2104	OE1	GLU	B	16	10803	15815	14286	-1237	1344	-3855 O
ATOM	2105	OE2	GLU	B	16	100.011	7.901	7.092	1.00101.66	O	
ANISOU	2105	OE2	GLU	B	16	9701	15696	13228	-1058	1471	-1847 O
ATOM	2106	N	PRO	B	17	96.750	6.893	8.820	1.00	60.06	N
ANISOU	2106	N	PRO	B	17	8523	19626	11691	-748	1094	-1600 N

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ATOM	2107	CA	PRO	B	17	37.354	5.868	9-737	1.00	87.03	C
ANISOU	2107	CA	PRO	B	17	8373	13551	11453	-570	965	-1595 C
ATOM	2108	C	PRO	B	17	96.738	4.436	3.865	1.00	68.46	C
ANISOU	2108	C	PRO	B	17	8306	13631	11637	-291	870	-1507 C
ATOM	2109	O	PRO	B	17	37.320	3.634	10.539	1.00	86.28	O
ANISOU	2109	O	PRO	B	17	7862	19508	11413	-144	773	-1488 O
ATOM	2110	CB	PRO	B	17	97.368	6.563	31.113	1.00	81.42	C
ANISOU	2110	CB	PRO	B	17	7331	12922	10631	-710	891	-1512 C
ATOM	2111	CG	PRO	B	17	97.098	8.017	30.833	1.00	86.45	C
ANISOU	2111	CG	PRO	B	17	3303	13455	11259	-989	1010	-1662 C
ATOM	2112	CD	PRO	B	17	36.231	3.0313	9.602	1.00	55.64	C
ANISOU	2112	CD	PRO	B	17	8211	13106	11224	-943	1101	-1609 C
ATOM	2113	N	PHE	B	18	95.610	4.104	9.185	1.00	84.80	N
ANISOU	2113	N	PHE	B	18	8054	12917	11259	-234	835	-1453 N
ATOM	2114	CA	PHE	B	18	94.928	2.759	9.317	1.00	33.29	C
ANISOU	2114	CA	PHE	B	18	7934	12691	11121	28	818	-1876 C
ATOM	2115	C	PHE	B	18	95.580	1.495	6.739	1.00	82.63	C
ANISOU	2115	C	PHE	B	18	7744	12613	11108	244	823	-1300 C
ATOM	2116	O	PHE	B	18	95.313	1.528	7.736	1.00	87.06	O
ANISOU	2116	O	PHE	B	18	8328	13334	11757	228	927	-1452 O
ATOM	2117	CB	PHE	B	18	93.435	2.743	8.722	1.00	76.40	C
ANISOU	2117	CB	PHE	B	18	7333	11444	10271	93	340	-3326 C
ATOM	2118	CG	PHE	B	18	92.541	3.794	9.282	3.00	77.63	C
ANISOU	2118	CG	PHE	B	18	7619	11475	10402	-122	830	-1295 C
ATOM	2119	CD1	PHE	B	18	92.662	4.274	10.536	1.00	79.88	C
ANISOU	2119	CD1	PHE	B	18	7859	11835	10357	-107	763	-1290 C
ATOM	2120	CD2	PHE	B	18	91.490	4.286	8.491	1.00	74.53	C
ANISOU	2120	CD2	PHE	B	18	7423	10303	10015	-183	336	-1267 C
ATOM	2121	CE1	PHE	B	18	91.776	5.278	11.070	1.00	78.16	C
ANISOU	2121	CE1	PHE	B	18	7790	11486	10420	-333	776	-1268 C
ATOM	2122	CE2	PHE	B	18	30.607	5.246	8.079	1.00	76.37	C
ANISOU	2122	CE2	PHE	B	18	7786	11004	10223	-302	339	-1230 C
ATOM	2123	CZ	PHE	B	18	93.754	5.729	10.259	1.00	75.98	C
ANISOU	2123	CZ	PHE	B	18	7600	11012	10150	-973	843	-1235 C
ATOM	2124	N	SER	B	19	95.572	0.302	9.545	1.30	79.21	N
ANISOU	2124	N	SER	B	19	7203	12145	10690	449	737	-1825 N
ATOM	2125	CA	SER	B	19	95.007	-0.977	9.180	1.00	73.03	C
ANISOU	2125	CA	SER	B	19	6526	11482	10082	636	746	-1813 C
ATOM	2126	C	SER	B	19	95.188	-1.500	7.999	1.00	74.48	C
ANISOU	2126	C	SER	B	19	6784	11324	10190	743	824	-1328 C

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ATOM	2127	O	SER	B	19	34.135	-3.911	7.667	1.00	72.01	O	
ANISOU	2127	O	SER	B	19	6602	10587	9805	611	838	-1320	O
ATOM	2128	CB	SER	B	19	95.724	-1.300	10.364	1.00	70.53	C	
ANISOU	2128	CB	SER	B	19	6105	11016	9679	378	691	-1214	C
ATOM	2129	OG	SER	B	19	94.298	-2.037	13.557	1.30	60.33	O	
ANISOU	2129	OG	SER	B	19	5051	9467	3403	850	602	-1158	O
ATOM	2130	N	LYS	B	20	95.696	-2.603	7.337	1.00	69.84	N	
ANISOU	2130	N	LYS	B	20	6090	10665	9591	930	876	-1350	N
ATOM	2131	CA	LYS	B	20	94.952	-3.154	6.199	1.00	58.26	C	
ANISOU	2131	CA	LYS	B	20	6120	10335	9477	964	953	-1387	C
ATOM	2132	C	LYS	B	20	93.465	-3.446	5.405	1.00	67.11	C	
ANISOU	2132	C	LYS	B	20	6258	9991	9403	956	906	-1331	C
ATOM	2133	O	LYS	B	20	92.654	-6.056	5.570	1.00	62.77	O	
ANISOU	2133	O	LYS	B	20	5804	9275	8771	842	946	-1358	O
ATOM	2134	CB	LYS	B	20	95.574	-4.384	5.641	1.00	67.73	C	
ANISOU	2134	CB	LYS	B	20	5982	10282	9459	1180	1030	-1426	C
ATOM	2135	CG	LYS	B	20	97.039	-4.075	5.051	1.00	72.60	C	
ANISOU	2135	CG	LYS	B	20	6384	11132	10067	1171	1118	-1503	C
ATOM	2136	CD	LYS	B	20	97.525	-5.200	4.142	1.00	75.92	C	
ANISOU	2136	CD	LYS	B	20	6787	11517	10541	1358	1229	-1564	C
ATOM	2137	CE	LYS	B	20	98.641	-4.109	3.237	1.00	73.15	C	
ANISOU	2137	CE	LYS	B	20	6902	12006	10790	1292	1347	-1560	C
ATOM	2138	NZ	LYS	B	20	99.084	-5.729	2.229	1.00	81.54	N	
ANISOU	2138	NZ	LYS	B	20	7304	12665	11535	1455	1480	-1737	N
ATOM	2139	N	LYS	B	21	93.112	-4.130	7.498	1.00	67.70	N	
ANISOU	2139	N	LYS	B	21	6309	9945	9567	1079	820	-1250	N
ATOM	2140	CA	LYS	B	21	91.704	-4.476	7.765	1.00	56.03	C	
ANISOU	2140	CA	LYS	B	21	6304	9506	9216	1010	777	-1195	C
ATOM	2141	C	LYS	B	21	90.839	-3.280	7.074	1.00	54.71	C	
ANISOU	2141	C	LYS	B	21	6225	9809	9058	863	740	-1177	C
ATOM	2142	O	LYS	B	21	89.758	-6.157	7.434	1.30	65.33	O	
ANISOU	2142	O	LYS	B	21	6422	9192	9096	795	755	-1181	O
ATOM	2143	CB	LYS	B	21	91.534	-5.417	5.948	1.00	69.94	C	
ANISOU	2143	CB	LYS	B	21	6815	9935	9824	1235	704	-1106	C
ATOM	2144	CG	LYS	B	21	91.483	-6.593	5.586	1.00	50.03	C	
ANISOU	2144	CG	LYS	B	21	8163	11065	11179	1455	756	-1102	C
ATOM	2145	CD	LYS	B	21	90.950	-7.112	9.765	1.00	88.75	C	
ANISOU	2145	CD	LYS	B	21	9346	12048	12327	1560	539	-999	C
ATOM	2146	CE	LYS	B	21	91.836	-7.565	11.010	1.00	89.16	C	
ANISOU	2146	CE	LYS	B	21	9231	12293	12351	1549	533	-914	C

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ATOM	2147	NZ	LYS	B	21	91.107	-6.026	12.224	1.00	92.50	N
ANISOU	2147	NZ	LYS	B	21	9758	12604	12784	1733	528	-798 N
ATOM	2148	N	LEU	B	22	91.380	-2.249	8.739	1.00	61.18	N
ANISOU	2148	N	LEU	B	22	5674	9007	8565	751	700	-1162 N
ATOM	2149	CA	LEU	B	22	90.595	-0.900	3.942	1.00	56.73	C
ANISOU	2149	CA	LEU	B	22	5196	8404	7955	556	568	-1146 C
ATOM	2150	C	LEU	B	22	93.443	-0.242	7.634	1.00	54.52	C
ANISOU	2150	C	LEU	B	22	4975	8101	7640	437	777	-1203 C
ATOM	2151	O	LEU	B	22	89.408	6.343	7.275	1.00	52.07	O
ANISOU	2151	O	LEU	B	22	4804	7668	7213	341	782	-1178 O
ATOM	2152	CB	LEU	B	22	91.291	-0.091	9.989	1.00	60.57	C
ANISOU	2152	CB	LEU	B	22	5557	9062	8393	468	645	-1144 C
ATOM	2153	CG	LEU	B	22	90.360	-0.153	11.457	1.00	67.54	C
ANISOU	2153	CG	LEU	B	22	6501	9969	9308	490	545	-1072 C
ATOM	2154	CD1	LEU	B	22	90.867	-1.691	11.964	1.00	68.31	C
ANISOU	2154	CC1	LEU	B	22	6552	9987	9415	711	493	-1010 C
ATOM	2155	CD2	LEU	B	22	91.748	0.597	12.342	1.00	71.94	C
ANISOU	2155	CD2	LEU	B	22	6874	10694	9762	330	514	-1095 C
ATOM	2156	N	ARG	B	23	91.485	-0.245	6.805	1.00	52.95	N
ANISOU	2156	N	ARG	B	23	4655	5032	7425	441	852	-1271 N
ATOM	2157	CA	ARG	B	23	91.4153	0.492	5.545	1.00	53.40	C
ANISOU	2157	CA	ARG	B	23	4775	3382	7432	315	945	-1317 C
ATOM	2158	C	ARG	B	23	90.458	-0.210	4.574	1.00	51.02	C
ANISOU	2158	C	ARG	B	23	4530	7517	7137	372	954	-1320 C
ATOM	2159	O	ARG	B	23	89.526	0.447	3.755	1.00	52.47	O
ANISOU	2159	O	ARG	B	23	4923	7742	7270	266	1002	-1317 O
ATOM	2160	CB	ARG	B	23	92.874	0.683	4.930	1.00	57.54	C
ANISOU	2160	CB	ARG	B	23	5127	8804	7931	292	1034	-1396 C
ATOM	2161	CG	ARG	B	23	92.987	1.455	3.601	1.00	60.85	C
ANISOU	2161	CG	ARG	B	23	5596	9238	8286	158	1148	-1442 C
ATOM	2162	CD	ARG	B	23	92.496	2.931	3.625	1.00	61.20	C
ANISOU	2162	CD	ARG	B	23	5737	9241	8277	-46	1172	-1406 C
ATOM	2163	NE	ARG	B	23	92.393	3.528	2.257	1.00	62.96	N
ANISOU	2163	NE	ARG	B	23	6050	9441	8432	-145	1280	-1423 N
ATOM	2164	CZ	ARG	B	23	93.387	4.146	1.596	1.00	56.75	C
ANISOU	2164	CZ	ARG	B	23	6446	10051	8864	-251	1392	-1477 C
ATOM	2165	NH1	ARG	B	23	94.579	4.287	2.161	1.00	66.11	N
ANISOU	2165	NH1	ARG	B	23	6171	1.146	8801	-284	1409	-1530 N
ATOM	2166	NH2	ARG	B	23	93.201	4.631	0.369	1.00	59.61	N
ANISOU	2166	NH2	ARG	B	23	5846	9116	7888	-330	1487	-1477 N

TABLE 3-continued

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ATOM	2167	N	VAL	B	24	90.322	-1.528	4.661	1.00	48.90	N
ANISOU	2167	N	VAL	B	24	4377	7276	5925	535	941	-1325 N
ATOM	2168	CA	VAL	B	24	89.298	-2.219	3.857	1.00	50.40	C
ANISOU	2168	CA	VAL	B	24	4722	7307	7119	565	952	-1340 C
ATOM	2169	C	VAL	B	24	87.888	-1.651	4.196	1.00	46.68	C
ANISOU	2169	C	VAL	B	24	4390	6711	6636	471	887	-1270 C
ATOM	2170	O	VAL	B	24	87.091	-1.286	3.286	1.00	46.01	O
ANISOU	2170	O	VAL	B	24	4410	6570	6501	389	907	-1274 O
ATOM	2171	CB	VAL	B	24	89.334	-3.746	4.100	1.00	50.43	C
ANISOU	2171	CB	VAL	B	24	4737	7225	7199	748	944	-1354 C
ATOM	2172	CG1	VAL	B	24	88.113	-4.423	3.474	1.00	51.21	C
ANISOU	2172	CG1	VAL	B	24	5007	7148	7303	745	944	-1373 C
ATOM	2173	CG2	VAL	B	24	90.638	-4.384	3.564	1.00	52.95	C
ANISOU	2173	CG2	VAL	B	24	4932	7651	7535	864	1029	-1428 C
ATOM	2174	N	VAL	B	25	87.601	-1.551	5.490	1.00	48.29	N
ANISOU	2174	N	VAL	B	25	4584	5886	5878	486	813	-1203 N
ATOM	2175	CA	VAL	B	25	86.272	-1.020	5.917	1.00	48.89	C
ANISOU	2175	CA	VAL	B	25	4778	6846	5953	408	760	-1136 C
ATOM	2176	C	VAL	B	25	86.096	0.464	5.618	1.00	51.60	C
ANISOU	2176	C	VAL	B	25	5145	7225	7235	256	790	-1115 C
ATOM	2177	O	VAL	B	25	85.067	0.867	5.010	1.00	50.71	O
ANISOU	2177	O	VAL	B	25	5140	7032	7095	198	792	-1086 O
ATOM	2178	CB	VAL	B	25	85.745	-1.555	7.294	1.00	52.24	C
ANISOU	2178	CB	VAL	B	25	5224	7190	7434	477	688	-1071 C
ATOM	2179	CG1	VAL	B	25	86.812	-1.879	8.279	1.00	49.75	C
ANISOU	2179	CG1	VAL	B	25	4786	6979	7139	562	657	-1061 C
ATOM	2180	CG2	VAL	B	25	84.576	-0.723	7.912	1.00	46.41	C
ANISOU	2180	CG2	VAL	B	25	4574	6371	6890	382	643	-1004 C
ATOM	2181	N	ALA	B	26	87.137	1.246	5.922	1.00	45.34	N
ANISOU	2181	N	ALA	B	26	4251	6460	6418	192	822	-1132 N
ATOM	2182	CA	ALA	B	26	87.180	2.668	5.563	1.00	45.19	C
ANISOU	2182	CA	ALA	B	26	4258	6589	6343	40	880	-1122 C
ATOM	2183	C	ALA	B	26	86.920	2.859	4.056	1.00	45.79	C
ANISOU	2183	C	ALA	B	26	4408	6628	6364	5	945	-1137 C
ATOM	2184	O	ALA	B	26	86.073	3.639	3.671	1.00	44.41	O
ANISOU	2184	O	ALA	B	26	4339	6379	6154	-65	958	-1084 O
ATOM	2185	CB	ALA	B	26	86.552	3.276	5.947	1.00	45.27	C
ANISOU	2185	CB	ALA	B	26	4127	6741	6333	-31	921	-1169 C
ATOM	2186	N	ASP	B	27	87.705	2.200	3.200	1.00	44.99	N
ANISOU	2186	N	ASP	B	27	4246	6604	6244	54	994	-1207 N

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ATOM	2187	CA	ASP	B	27	87.436	2.166	1.752	1.00	45.42	C
ANISOU	2187	CA	ASP	B	27	4376	6652	6231	33	1050	-1230 C
ATOM	2188	C	ASP	B	27	86.045	1.708	1.349	1.00	45.94	C
ANISOU	2188	C	ASP	B	27	4571	6597	6289	65	994	-1195 C
ATOM	2189	O	ASP	B	27	85.439	2.257	0.419	1.00	43.73	O
ANISOU	2189	O	ASP	B	27	4379	6304	5933	3	1016	-1166 O
ATOM	2190	CB	ASP	B	27	86.421	1.230	1.020	1.00	47.73	C
ANISOU	2190	CB	ASP	B	27	4586	7031	6517	109	1109	-1325 C
ATOM	2191	CG	ASP	B	27	89.801	1.854	0.827	1.00	52.63	C
ANISOU	2191	CG	ASP	B	27	5079	7807	7110	44	1199	-1371 C
ATOM	2192	OD1	ASP	B	27	89.930	3.101	0.925	1.00	56.56	O
ANISOU	2192	OD1	ASP	B	27	5585	6335	7572	-90	1235	-1336 O
ATOM	2193	OD2	ASP	B	27	90.749	1.077	0.601	1.00	51.71	O
ANISOU	2193	OD2	ASP	B	27	4853	7779	7014	129	1241	-1445 O
ATOM	2194	N	TYR	B	28	85.558	0.686	2.007	1.00	45.07	N
ANISOU	2194	N	TYR	B	28	4468	6408	6249	160	925	-1199 N
ATOM	2195	CA	TYR	B	28	84.178	0.204	1.727	1.00	46.87	C
ANISOU	2195	CA	TYR	B	28	4805	6529	6475	172	868	-1174 C
ATOM	2196	C	TYR	B	28	83.111	1.284	1	999	1.00	45.62 C
ANISOU	2196	C	TYR	B	28	4716	6319	6299	98	832	-1075 C
ATOM	2197	O	TYR	B	28	82.213	1.490	1.202	1.00	42.86	O
ANISOU	2197	O	TYR	B	28	4440	5950	5893	86	818	-1047 O
ATOM	2198	CB	TYR	B	28	83.833	6897	6824	273	812	-1187 C
ANISOU	2198	CB	TYR	B	28	5067	6697	6824	273	812	-1187 C
ATOM	2199	CG	TYR	B	28	82.455	-1.589	2.226	1.00	47.10	C
ANISOU	2199	CG	TYR	B	28	4936	6367	6593	265	764	-1180 C
ATOM	2200	CD1	TYR	B	28	82.282	-2.555	1.209	1	00	45.86 C
ANISOU	2200	CD1	TYR	B	28	4826	6194	6406	282	787	-1266 C
ATOM	2201	CD2	TYR	B	28	81.332	-1.142	2.918	1.00	43.19	C
ANISOU	2201	CD2	TYR	B	28	4484	5804	6123	231	702	-1097 C
ATOM	2202	CE1	TYR	B	28	81.025	-3.058	0.838	1.00	48.29	C
ANISOU	2202	CE1	TYR	B	28	5212	6431	6705	251	742	-1274 C
ATOM	2203	CE2	TYR	B	28	80.063	-1.637	2.598	1.00	40.85	C
ANISOU	2203	CE2	TYR	B	28	4257	5442	5825	213	657	-1096 C
ATOM	2204	CZ	TYR	B	28	79.905	-2.583	1.601	1.00	44.29	C
ANISOU	2204	CZ	TYR	B	28	4729	5872	6227	216	672	-1184 C
ATOM	2205	OH	TYR	B	28	78.626	-3.090	1.281	1.00	45.73	O
ANISOU	2205	OH	TYR	B	28	4969	6006	6401	176	624	-1196 O
ATOM	2206	N	ILE	B	29	83.239	1.968	3.140	1.00	44.02	N
ANISOU	2206	N	ILE	B	29	4484	6101	6140	75	818	-1024 N

TABLE 3-continued

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ATOM	2207	CA	ILE	B	29	82.281	2.979	3.530	1.00	42.77	C
ANISOU	2207	CA	ILE	B	29	4392	5878	5960	20	800	-935 C
ATOM	2208	C	ILE	B	29	82.361	4.140	2.539	1.00	44.51	C
ANISOU	2208	C	ILE	B	29	4659	6135	6118	-61	869	-900 C
ATOM	2209	O	ILE	B	29	81.319	4.595	2.094	1.00	41.92	O
ANISOU	2209	O	ILE	B	29	4408	5763	5758	-71	853	-830 O
ATOM	2210	CB	ILE	B	29	82.540	3.398	4.979	1.00	45.76	C
ANISOU	2210	CB	ILE	B	29	4734	6238	6416	F6	786	-910 C
ATOM	2211	CG1	ILE	B	29	82.392	2.207	5.916	1.00	41.82	C
ANISOU	2211	CG1	ILE	B	29	4204	5699	5985	97	718	-925 C
ATOM	2212	CG2	ILE	B	29	81.744	4.615	5.402	1.00	47.54	C
ANISOU	2212	CG2	ILE	B	29	5030	6394	6640	-58	799	-830 C
ATOM	2213	CD1	ILE	B	29	82.938	2.545	7.287	1.00	47.50	C
ANISOU	2213	CD1	ILE	B	29	4869	6444	6733	84	704	-913 C
ATOM	2214	N	LEU	B	30	83.580	4.571	2.129	1.00	41.39	N
ANISOU	2214	N	LEU	B	30	4215	5830	5683	-113	949	-944 N
ATOM	2215	CA	LEU	B	30	83.733	5.661	1.163	1.00	45.54	C
ANISOU	2215	CA	LEU	B	30	4797	6383	6124	-196	1032	-906 C
ATOM	2216	C	LEU	B	30	83.001	5.385	-0.158	1.00	45.89	C
ANISOU	2216	C	LEU	B	30	4914	6439	6085	-172	1020	-685 C
ATOM	2217	O	LEU	B	30	82.340	6.281	-0.668	1.00	45.60	O
ANISOU	2217	O	LEU	B	30	4962	6374	5991	-203	1039	-793 O
ATOM	2218	CB	LEU	B	30	85.201	6.017	0.860	1.00	40.30	C
ANISOU	2218	CB	LEU	B	30	4438	6205	5808	-265	1130	-972 C
ATOM	2219	CG	LEU	B	30	35.774	6.370	2.007	1.00	44.63	C
ANISOU	2219	CG	LEU	B	30	4562	8371	8024	-345	1160	-970 C
ATOM	2220	CD1	LEU	B	30	87.303	68.30	2.022	1.00	45.29	C
ANISOU	2220	CD1	LEU	B	30	4512	6094	8102	-304	1222	-1062 C
ATOM	2221	CD2	LEU	B	30	85.287	8.317	1.960	1.00	45	45 C
ANISOU	2221	CD2	LEU	B	30	4781	5386	6101	-439	1226	-862 C
ATOM	2222	N	GLU	B	31	63.142	4.160	-0.584	1.00	45.41	N
ANISOU	2222	N	GLU	B	31	4943	5547	5138	-115	903	-968 N
ATOM	2223	CA	GLU	B	31	82.600	3.804	-1.980	1.00	47.74	C
ANISOU	2223	CA	GLU	B	31	5184	5760	5215	-110	985	-977 C
ATOM	2224	C	GLU	B	31	81.159	3.403	-1.375	1.00	44.12	C
ANISOU	2224	C	GLU	B	31	4787	S235	8783	-73	884	-933 C
ATOM	2225	O	GLU	B	31	80.485	3.280	-2.856	1.00	41.60	O
ANISOU	2225	O	GLU	B	31	4500	5957	5350	-84	861	-912 O
ATOM	2226	CB	GLU	B	31	33.417	2.093	-2.598	1.00	49.97	C
ANISOU	2226	CB	GLU	B	31	5412	7101	6471	-78	1019	-1104 C

TABLE 3-continued

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ATOM	2227	CG	GLU	B	31	84.799	3.286	-2.989	1.00	56.54	C	
ANISOU	2227	CG	GLU	B	31	5192	8026	7263	-129	1134	-1140	C
.ATOM	2228	CD	GLU	B	31	85.913	2.231	-3.059	1.00	62.88	C	
ANISOU	2228	CD	GLU	B	31	8902	8890	8130	-74	1131	-1255	O
ATOM	2229	OE1	GLU	B	31	35.621	1.012	-3.233	1.00	61	97	O
ANISOU	2229	OE1	GLU	B	31	6795	8744	8307	1	3146	-1336	O
ATOM	2230	OE2	GLU	B	31	87.092	2.672	-3.007	1.00	67.07	O	
ANISOU	2230	OE2	GLU	B	31	7349	9502	3633	-310	1254	-1294	O
ATOM	2231	N	ASN	B	32	50.705	3.033	-0.679	1.00	42.67	N	
ANISOU	2231	N	ASN	B	32	4557	5968	5588	-34	823	-922	N
ATOM	2232	CA	ASN	B	32	79.293	2.674	-0.529	1.30	42.38	C	
ANISOU	2232	CA	ASN	B	32	4554	5877	5670	-10	735	-830	C
ATOM	2233	C	ASN	B	32	78.507	3.491	0.503	1.00	41.87	C	
ANISOU	2233	C	ASN	B	32	4502	5734	5072	-6	706	-774	C
ATOM	2234	O	ASN	B	32	77.536	2.305	1.028	1.00	42.57	O	
ANISOU	2234	O	ASN	B	32	4539	5771	5815	22	639	-751	O
ATOM	2235	CB	ASN	B	32	79.119	1.185	-0.207	1.00	39.37	C	
ANISOU	2235	CB	ASN	B	32	4152	5455	5352	34	889	-973	C
ATOM	2236	CB	ASN	B	32	72.386	0.260	-1.155	1.00	44.49	C	
ANISOU	2236	CG	ASN	B	32	4796	5159	5949	41	733	-1091	C
ATOM	2237	CG	ASN	B	32	81.029	-0.120	-0.361	1.00	40.90	O	
ANISOU	2237	OD1	ASN	B	32	5053	6471	6297	78	788	-154	O
ATOM	2238	OD1	ASN	B	32	79.304	0.011	-2.011	1.00	40.21	N	
ANISOU	2238	ND1	ASN	B	32	4306	5873	5314	8	710	-1123	N
ATOM	2239	N	ALA	B	33	79.893	4.768	0.730	1.30	43.80	N	
ANISOU	2239	N	ALA	B	33	4387	5587	5529	-41	769	-705	N
ATOM	2240	CA	ALA	B	33	78.300	5.6134	1.792	3.30	39.74	C	
ANISOU	2240	OA	ALA	B	33	4273	5208	5401	-42	771	-021	C
ATOM	2241	C	ALA	B	33	75.802	5.583	1.877	1.00	38.74	C	
ANISOU	2241	C	ALA	B	33	4169	5199	5353	700	-544	C	
ATOM	2242	O	ALA	B	33	76.210	5.415	2.972	1.00	39.50	O	
ANISOU	2242	O	ALA	B	33	4251	5224	5535	24	657	-529	O
ATOM	2243	CB	ALA	B	33	78.347	7.077	1.597	1.00	38.04	C	
ANISOU	2243	CB	ALA	B	33	4173	5203	5270	-99	869	-557	C
ATOM	2244	N	HIS	B	34	75.154	5.693	0.724	1.00	39.88	N	
ANISOU	2244	N	HIS	B	34	4333	5405	5409	11	674	-500	N
ATOM	2245	CA	HIS	B	34	74.894	5.825	0.698	1.00	42.59	C	
ANISOU	2245	CA	HIS	B	34	4083	5742	5757	52	608	-414	C
ATOM	2246	C	HIS	B	34	74.074	4.345	1.152	1.00	42.33	O	
ANISOU	2246	C	HIS	B	34	4568	5009	57115	62	529	-486	C

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TABLE 3-continued

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ATOM	2247	O	HIS	B	34	73.140	4.570	1.955	1.00	43.74	O
ANISOU	2247	O	HIS	B	34	4750	5825	6036	87	496	-440
ATOM	2248	CB	HIS	B	34	74.146	5.221	-0.689	1.00	46.83	C
ANISOU	2248	CB	HIS	B	34	5243	6375	6163	64	585	-342
ATOM	2249	CG	HIS	B	34	72.640	3.480	-0.691	1.00	48.79	C
ANISOU	2249	CG	HIS	B	34	5473	6420	119	513	-238	C
ATOM	2250	ND1	HIS	B	34	71.732	5.490	-0.694	1.00	49.43	N
ANISOU	2250	ND1	HIS	B	34	5492	6776	5514	119	421	-284
ATOM	2251	CD2	HIS	B	34	71.910	7.872	-0.545	1.00	52.33	C
ANISOU	2251	CD2	HIS	B	34	5953	7073	6872	180	539	-85
ATOM	2252	CE1	HIS	B	34	73.483	6.305	-0.683	1.00	44.93	C
ANISOU	2252	CE1	HIS	B	34	4890	6234	5947	174	375	-170
ATOM	2253	NE2	HIS	B	34	70.585	7.335	-0.846	1.00	49.63	N
ANISOU	2253	NE2	HIS	B	34	5543	6779	6537	223	447	-43
ATOM	2254	N	ASP	B	35	74.591	3.406	0.698	1.00	44.46	N
ANISOU	2254	N	ASP	B	35	4853	6010	6030	42	511	-602
ATOM	2255	CA	ASP	B	35	74.019	2.121	1.355	1.00	40.24	C
ANISOU	2255	CA	ASP	B	35	4377	5529	5648	43	452	-675
ATOM	2256	C	ASP	B	35	74.257	1.754	2.524	1.00	40.97	C
ANISOU	2256	C	ASP	B	35	4368	5424	5773	65	468	-695
ATOM	2257	O	ASP	B	35	73.571	1.081	3.308	1.00	39.32	O
ANISOU	2257	O	ASP	B	35	4147	5160	5534	70	425	-702
ATOM	2258	CB	ASP	B	35	74.372	0.994	0.182	1.00	50.37	C
ANISOU	2258	CB	ASP	B	35	5577	6780	5780	16	442	-797
ATOM	2259	CG	ASP	B	35	73.463	-0.189	0.348	1.00	59.21	C
ANISOU	2259	CG	ASP	B	35	6684	7871	7942	-5	384	-851
ATOM	2260	OD1	ASP	B	35	73.977	-1.312	0.496	1.00	61.43	O
ANISOU	2260	OD1	ASP	B	35	6979	8097	8254	-6	405	-968
ATOM	2261	OD2	ASP	B	35	72.220	0.013	0.404	1.00	65.29	O
ANISOU	2261	OD2	ASP	B	35	7429	8666	8714	-18	325	-502
ATOM	2262	N	VAL	B	36	75.492	2.219	3.128	1.00	37.49	N
ANISOU	2262	N	VAL	B	36	3928	4964	5351	71	526	-599
ATOM	2263	CA	VAL	B	36	75.933	1	815	4.457	1.00	36.59
ANISOU	2263	CA	VAL	B	36	3797	4778	5328	93	532	-720
ATOM	2264	C	VAL	B	36	74.955	2.404	5.509	1.00	26.52	C
ANISOU	2264	C	VAL	B	36	3070	4075	5253	97	519	-636
ATOM	2265	O	VAL	B	36	74.850	1.332	6.598	1.00	35.98	CO
ANISOU	2265	O	VAL	B	36	3722	4571	5379	115	505	-645
ATOM	2266	CB	VAL	B	36	77.395	2.251	4.728	1.00	39.46	C
ANISOU	2266	CB	VAL	B	36	4138	5173	5682	37	590	-747

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TABLE 3-continued

ATOM	2267	CG1	VAL	B	36	77.362	2.413	6.221	1.00	37.15	C
ANISOU	2267	CG1	VAL	B	36	3829	4830	5458	97	593	-728
ATOM	2268	CG2	VAL	B	36	78.392	1.287	4.028	1.00	40.39	C
ANISOU	2268	CG2	VAL	B	36	4228	5343	5774	109	606	-847
ATOM	2269	N	GLN	B	37	74.247	3.499	5.150	1.00	37.58	N
ANISOU	2269	N	GLN	B	37	3950	4845	5482	89	530	-550
ATOM	2270	CA	GLN	B	37	73.217	4.132	6.057	1.00	43.05	C
ANISOU	2270	CA	GLN	B	37	4650	5473	6233	105	532	-468
ATOM	2271	C	GLN	B	37	72.223	3.008	6.505	1.00	44.25	C
ANISOU	2271	C	GLN	B	37	4774	5598	6441	115	474	-486
ATOM	2272	O	GLN	B	37	71.683	3.135	7.615	1.00	40.05	O
ANISOU	2272	O	GLN	B	37	4242	5000	5975	124	482	-455
ATOM	2273	CB	GLN	B	37	72.322	5.112	5.287	1.00	40.78	C
ANISOU	2273	CB	GLN	B	37	4375	5215	5933	124	533	-370
ATOM	2274	CG	GLN	B	37	72.967	5.365	4.874	1.33	49.28	C
ANISOU	2274	CG	GLN	B	37	5502	6290	5931	115	610	-319
ATOM	2275	CD	GLN	B	37	71.955	7.137	4.140	1.00	44.63	C
ANISOU	2275	CD	GLN	B	37	4926	5727	9304	160	905	-208
ATOM	2276	OE1	GLN	B	37	71.781	7.904	2.897	1.00	44.17	O
ANISOU	2276	OE1	GLN	B	37	4860	5765	6155	168	567	-191
ATOM	2277	NE2	GLN	B	37	71.180	7.852	4.395	1.00	46.82	N
ANISOU	2277	NE2	GLN	B	37	5215	5930	6643	199	637	-129
ATOM	2278	N	PHE	B	38	71.955	2.121	5.601	1.00	38.24	N
ANISOU	2278	N	PHE	B	38	3995	4839	5647	102	425	-541
ATOM	2279	CA	PHE	B	38	70.925	1.118	5.817	1.00	39.79	C
ANISOU	2279	CA	PHE	B	38	4166	6065	5387	88	377	-568
ATOM	2280	C	PHE	B	38	71.446	-0.234	6.281	1.00	40.72	C
ANISOU	2280	C	PHE	B	38	4531	5119	6050	81	373	-553
ATOM	2281	O	PHE	B	38	70.545	-1.156	6.435	1.00	38.93	O
ANISOU	2281	O	PHE	B	38	4068	4360	5353	54	353	-690
ATOM	2282	CB	PHE	B	38	70.044	0.950	4.563	1.00	42.89	C
ANISOU	2282	CB	PHE	B	38	4526	5550	6212	61	322	-572
ATOM	2283	CG	PHE	B	38	69.506	2.237	4.055	1.00	42.03	C
ANISOU	2283	CG	PHE	B	38	4521	5638	6173	92	319	-453
ATOM	2284	CD1	PHE	B	38	68.447	2.856	4.726	1.00	43.53	C
ANISOU	2284	CD1	PHE	B	38	4555	5636	5266	124	319	-371
ATOM	2285	CD2	PHE	B	38	70.136	2.895	8.028	1.00	44.39	C
ANISOU	2285	CD2	PHE	B	38	4802	8962	6331	59	334	-442
ATOM	2286	CE1	PHE	B	38	67.996	4.091	4.283	1.00	48.05	C
ANISOU	2286	CE1	PHE	B	38	5121	6303	6831	177	330	-256

TABLE 3-continued

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ATOM	2287	CE2	PHL	B	38	69.687	4.127	2.582	1.00	50.65	C
ANISOU	2287	CE2	PHE	B	38	5521	6723	7001	142	345	-323 C
ATOM	2288	CZ	PHE	B	38	68.605	4.730	3.205	1.00	47.65	C
ANISOU	2288	CZ	PHE	B	38	6110	6316	6682	159	342	-226 C
ATOM	2289	N	GLN	B	39	72.753	-0.322	6.517	1.00	37.07	N
ANISOU	2289	N	GLN	B	39	3858	4540	5586	107	413	-693 N
ATOM	2290	CA	GLN	B	39	73.350	-1.571	6.932	1.00	37.93	C
ANISOU	2290	CA	GLN	B	39	3986	4686	5740	132	422	-751 C
ATOM	2291	C	GLN	B	39	73.551	-1.505	8.464	1.05	37.34	C
ANISOU	2291	C	GLN	B	39	3913	4543	5728	164	435	-718 C
ATOM	2292	O	GLN	B	39	73.817	-0.581	9.134	1.09	40.94	O
ANISOU	2292	O	GLN	B	39	4366	5009	6130	157	452	-553 O
ATOM	2293	CB	GLN	B	39	74.735	-1.835	5.312	1.00	35.26	C
ANISOU	2293	CB	GLN	B	39	3546	4391	5360	159	450	-829 C
ATOM	2294	CG	GLN	B	39	74.705	-1.905	4.795	1.00	97.67	C
ANISOU	2294	CG	GLN	B	39	3954	4773	5584	125	447	-885 C
ATOM	2295	CD	GLN	B	33	76.092	-2.223	4.175	1.00	40.26	C
ANISOU	2295	CD	GLN	B	39	4278	5143	5876	156	492	-961 C
ATOM	2296	OE1	GLN	B	39	77.050	-2.643	4.858	1.60	37.44	O
ANISOU	2296	OE1	GLN	B	39	3906	4754	5565	214	520	-979 O
ATOM	2297	NE2	GLN	B	39	75.185	-2.056	2.878	1.00	42.55	N
ANISOU	2297	NE2	GLN	B	39	4574	5515	6077	121	501	-1001 N
ATOM	2298	N	THR	B	40	73.478	-2.809	8.963	1.00	37.93	N
ANISOU	2298	N	THR	B	40	4021	4535	5854	184	435	-748 N
ATOM	2299	CA	THR	B	40	79.706	-3.047	10.375	1.30	38.13	C
ANISOU	2299	CA	THR	B	40	4061	4494	5923	224	445	-705 C
ATOM	2300	C	THR	B	40	75.207	-3.266	10.555	1.00	38	92 C
ANISOU	2300	C	THR	B	40	4149	4628	6010	292	453	-728 C
ATOM	2301	O	THR	B	40	75.972	-3.402	9.547	1.00	37.01	O
ANISOU	2301	O	THR	B	40	9888	4439	5736	208	469	-788 O
ATOM	2302	CB	THR	B	40	72.933	-4.920	10.606	1.00	98.14	C
ANISOU	2302	CB	THR	B	40	3859	4136	5735	221	450	-716 C
ATOM	2303	OG1	THR	B	40	73.510	-5.409	10.040	1.00	134.37	O
ANISOU	2303	OG1	THR	B	40	2740	3955	5592	245	4355	-795 O
ATOM	2304	CG2	THR	B	40	71.433	-4.225	10.610	1.00	35.97	C
ANISOU	2304	CG2	THR	B	40	3829	4103	5734	141	437	-704 C
ATOM	2305	N	ILE	B	41	75.557	-3.321	11.810	1.00	37.54	N
ANISOU	2305	N	ILE	B	41	3988	4449	5864	336	457	-582 N
ATOM	2306	CA	ILE	B	41	77.099	-3.521	12.048	1.00	38.34	C
ANISOU	2306	CA	ILE	B	41	4039	4591	5935	415	459	-698 C

TABLE 3-continued

ATOM	2307	C	ILE	B	41	77.425	-4.998	11.484	1.00	40.96	C
ANISOU	2307	C	ILE	B	41	4490	4860	6303	466	477	-749 C
ATOM	2308	O	ILE	B	41	78.503	-5.159	10.926	1.00	37.11	O
ANISOU	2308	O	ILE	B	41	3859	4435	5796	539	492	-794 O
ATOM	2309	CB	ILE	B	41	77.484	-3.568	13.523	1.00	41.46	C
ANISOU	2309	CB	ILE	B	41	4428	4996	5328	455	443	-635 C
ATOM	2310	CG1	ILE	B	41	79.014	-3.575	13.703	1.00	42.42	C
ANISOU	2310	CG1	ILE	B	41	4475	5229	545	526	433	-649 C
ATOM	2311	CG2	ILE	B	41	75.768	-4.656	14.363	1.00	42.03	C
ANISOU	2311	CG2	ILE	B	41	4574	4945	8453	495	445	-588 C
ATOM	2312	CD1	ILE	B	41	79.267	-3.472	15.138	1.00	43.15	C
ANISOU	2312	CD1	ILE	B	41	4626	5422	5550	553	404	-583 C
ATOM	2313	N	THR	B	42	76.487	-5.944	11.590	1.00	45.42	N
ANISOU	2313	N	THR	B	42	4390	4654	5272	478	488	-742 N
ATOM	2314	CA	THR	B	42	76.574	-7.310	11.393	1.00	39.33	C
ANISOU	2314	CA	THR	B	42	4320	4433	8189	533	525	-805 C
ATOM	2315	C	THR	B	42	76.633	-7.252	9.583	1.00	41.78	C
ANISOU	2315	C	THR	B	42	4815	4795	6465	493	541	-902 C
ATOM	2316	O	THR	B	42	77.753	-7.900	9.035	1.00	39.07	O
ANISOU	2316	O	THR	B	42	4271	4448	4226	567	579	-960 O
ATOM	2317	CB	THR	B	42	75.504	-8.259	11.513	1.00	40.50	C
ANISOU	2317	CB	THR	B	42	4561	4428	6399	496	547	-793 C
ATOM	2318	OG1	THR	B	42	75.144	-7.989	12.873	1.00	40.78	O
ANISOU	2318	OG1	THR	B	42	4605	4443	5444	502	530	-695 O
ATOM	2319	CG2	THR	B	42	75.925	-9.829	11.391	1.00	39.59	C
ANISOU	2319	CG2	THR	B	42	4531	4159	8342	584	639	-832 C
ATOM	2320	N	ASP	B	43	76.034	-6.431	8.886	1	1.00	40.57 N
ANISOU	2320	N	ASP	B	43	4455	4714	5283	383	517	-917 N
ATOM	2321	CA	ASP	B	43	76.229	-6.287	7.418	1.00	45.51	C
ANISOU	2321	CA	ASP	B	43	5067	5412	6653	348	528	-1.001 C
ATOM	2322	C	ASP	B	43	77.583	-5.768	6.999	1.00	43.42	C
ANISOU	2322	C	ASP	B	43	4733	5241	4528	400	548	-1020 C
ATOM	2323	O	ASP	B	43	76.164	-5.252	5.035	1.00	43.52	O
ANISOU	2323	O	ASP	B	43	4747	5274	6516	421	567	-1103 O
ATOM	2324	CB	ASP	B	43	75.270	-5.322	6.780	1.00	45.07	C
ANISOU	2324	CB	ASP	B	43	4976	5422	6726	248	490	-984 C
ATOM	2325	CG	ASP	B	43	73.932	-5.649	5.570	1.00	50.29	C
ANISOU	2325	CG	ASP	B	43	5571	8030	7407	172	471	-1006 C
ATOM	2326	OD1	ASP	B	43	73.795	-7.063	5.257	1.00	47.59	O
ANISOU	2326	OD1	ASP	B	43	5295	5515	7005	160	5.00	-1385 O

TABLE 3-continued

ATOM	2327	OD2	ASP	B	43	73.303	-5.316	6.724	1.00	50.23	O	
ANISOU	2327	OD2	ASP	B	43	5531	6067	7387	121	434	-944	O
ATOM	2328	N	LEU	B	44	78.051	-4.742	7.711	1.00	45.34	N	
ANISOU	2328	N	LEU	B	44	4917	5555	6755	407	530	-952	N
ATOM	2329	CA	LEU	B	44	79.246	-4	080	7.312	1.00	41.46	C
ANISOU	2329	CA	LEU	B	44	4356	5130	6215	424	552	-971	C
ATOM	2330	C	LEU	B	44	80.391	-5.055	7.531	1.00	41.45	C	
ANISOU	2330	C	LEU	B	44	4328	5174	6249	540	582	-1006	C
ATOM	2331	O	LEU	B	44	81.274	-5.116	6.577	1.00	39.61	O	
ANISOU	2331	O	LEU	B	44	4062	5016	5993	565	823	-1070	O
ATOM	2332	CB	LEU	B	44	79.455	-2.810	8.114	1.00	25.64	C	
ANISOU	2332	CB	LEU	B	44	3701	4635	5584	367	534	-902	C
ATOM	2333	CG	LEU	B	44	63.527	-1.833	7.650	1.00	39.73	C	
ANISOU	2333	CG	LEU	B	44	4025	5153	5915	358	564	-921	C
ATOM	2334	CD1	LEU	B	44	80.549	-1.587	6.174	1.00	40.53	C	
ANISOU	2334	CD1	LEU	B	44	4144	5307	5966	313	599	-971	C
ATOM	2335	CD2	LEU	B	44	80.232	-0.527	5.373	1.00	43.25	C	
ANISOU	2335	CD2	LEU	B	44	4457	5622	6347	285	557	-858	C
ATOM	2336	N	ALA	B	45	80.413	-5.744	8.675	1.00	38.54	N	
ANISOU	2336	N	ALA	B	45	3975	4731	5938	613	556	-955	N
ATOM	2337	CA	ALA	B	45	81.458	-6.767	8.314	1.00	41.84	C	
ANISOU	2327	CA	ALA	B	45	4369	5135	6395	761	597	-570	C
ATOM	2338	C	ALA	B	45	81.423	-7.875	7.842	1.00	45.73	C	
ANISOU	2338	C	ALA	B	45	4925	5536	6913	796	560	-1064	C
ATOM	2339	O	ALA	B	45	82.457	-8.255	7.256	1.00	44.12	O	
ANISOU	2339	O	ALA	B	45	4676	5379	6757	878	711	-1121	O
ATOM	3340	CB	ALA	B	45	81.335	-7.325	10.323	1.00	39.83	C	
ANISOU	2340	CB	ALA	B	45	4140	4506	6186	841	568	-881	C
ATOM	2341	N	ARG	B	46	80.248	-3.400	7.551	1.05	43.39	N	
ANISOU	2341	N	ARG	B	46	4808	5193	6714	726	655	-1091	N
ATOM	2342	CA	ARG	B	46	80.103	-9.388	5.458	1.00	45.78	C	
ANISOU	2342	CA	ARG	B	46	5515	5542	7455	722	730	-1202	C
ATOM	2343	C	ARG	B	46	80.103	-8.871	6.458	1.00	51.03	C	
ANISOU	2343	C	ARG	B	46	5732	7121	7533	670	750	-1291	C
ATOM	2344	O	ARG	B	46	81.284	-9.579	4.343	1.00	45.48	O	
ANISOU	2344	O	ARG	B	46	6165	5530	5362	733	530	-1382	O
ATOM	2345	CB	ARG	B	46	76.568	-9.083	6.320	1.00	51.11	C	
ANISOU	2345	CB	ARG	B	46	5876	5885	7648	516	725	-1231	C
ATOM	2346	CG	ARG	B	46	78.192	-10.814	7.473	1.00	62.37	C	
ANISOU	2346	CG	ARG	B	46	7388	7149	9160	675	736	-1157	C

TABLE 3-continued

ATOM	2347	CD	ARG	B	46	77.636	-12.121	5.905	1.00	65.37	C	
ANISOU	2347	CD	ARG	B	46	8360	7830	10054	630	609	-1257	C
ATOM	2348	NE	ARG	B	46	77.020	-13.309	7.901	1.00	73.09	N	
ANISOU	2348	NE	ARG	B	46	5964	5160	10645	648	836	-1212	N
ATOM	2349	CZ	ARG	B	46	75.363	-14.106	7.503	1.00	75.51	C	
ANISOU	2349	CZ	ARG	B	46	5520	3422	11123	530	908	-1290	C
ATOM	2350	NH1	ARG	B	46	75.229	-14.523	6.349	1.00	77.73	N	
ANISOU	2350	NH1	ARG	B	46	9721	6574	11250	510	553	-1437	N
ATOM	2251	NH2	ARG	B	46	75.824	-14.871	8.572	1.00	76.76	N	
ANISOU	2351	NH2	ARG	B	46	3642	8290	11233	601	541	-1226	N
ATOM	2352	N	ASN	B	47	30.128	-7.562	4.758	1.00	43.48	N	
ANISOU	2352	N	ASN	B	47	4736	5275	5508	561	707	-1265	N
ATOM	2353	CA	ASN	B	47	80.423	-7.133	3.440	1.00	40.50	C	
ANISOU	2353	CA	ASN	B	47	5597	5773	6500	500	733	-1335	C
ATOM	2354	C	ASN	B	47	81.501	-6.774	3.314	1.00	47.12	C	
ANISOU	2354	C	ASN	B	47	5084	5955	5861	575	778	-1246	C
ATOM	2355	O	ASN	B	47	82.407	-6.717	2.220	1.00	46.70	O	
ANISOU	2355	O	ASN	B	47	5023	5977	5747	557	331	-1423	O
ATOM	2356	CB	ASN	B	47	75.546	-5.930	3.065	1.00	45.17	O	
ANISOU	2356	CB	ASN	B	47	4922	5563	5557	378	676	-1287	C
ATOM	2357	CG	ASN	B	47	78.768	-6.301	8.007	1.00	50.54	C	
ANISOU	2357	CG	ASN	B	47	5658	6290	7245	300	632	-1288	C
ATOM	2358	OD1	ASN	B	47	77.752	-7.404	2.614	1.00	46.13	O	
ANISOU	2358	OD1	ASN	B	47	5181	5665	6713	285	555	-1374	O
ATOM	2359	NH2	ASN	B	47	77.213	-5.356	3.445	1.00	42.75	N	
ANISOU	2359	NH2	ASN	B	47	4564	5330	3250	243	573	-1199	N
ATOM	2360	N	THR	B	48	32.587	-5.543	4.425	1.00	45.79	N	
ANISOU	2360	N	THR	B	48	4842	5817	5733	652	759	-1273	N
ATOM	2361	CA	THR	B	48	34.001	-5.186	4.217	1.00	46.85	C	
ANISOU	2361	CA	THR	B	48	4565	6085	6555	710	726	-1259	C
ATOM	2362	C	THR	B	48	54.008	-7.330	4.555	1.00	49.56	C	
ANISOU	2362	C	THR	B	48	5178	5387	7266	875	850	-1321	C
ATOM	2363	O	THR	B	48	88.149	-7.273	4.424	1.06	47.49	O	
ANISOU	2363	O	THR	B	48	4804	6242	6997	546	891	-1343	O
ATOM	2364	CB	THR	B	48	84.328	-5.021	5.236	1.00	47.70	C	
ANISOU	2364	CB	THR	B	48	4334	6292	6946	672	745	-1204	C
ATOM	0	2365	OG1	THR	B	48	84.144	-5.398	6.612	1.00	51.08	O
ANISOU	2365	OG1	THR	B	48	5311	6659	7435	741	695	-1127	O
ATOM	2366	CG2	THR	B	48	53.393	-3.811	4.901	1.90	46.15	C	
ANISOU	2366	CG2	THR	B	48	4733	6113	6588	523	718	-1159	C

TABLE 3-continued

ATOM	2367	N	GLN	B	49	54.259	-8.510	4.357	1.00	49.56	N
ANISOU	2367	N	GLN	B	49	5278	6219	7333	936	856	-1320 N
ATOM	2368	CA	GLN	B	49	84.982	-9.718	5.349	1.00	54.89	C
ANISOU	2368	CA	OLN	B	49	5954	6312	8039	1116	905	-1319 C
ATOM	2369	C	GLN	B	49	85.964	-9.382	6.484	1.30	58.64	C
ANISOU	2369	C	GLN	B	49	6235	7434	8582	1224	662	-1224 C
ATOM	2370	O	GLN	B	49	81.100	-9.812	8.432	1.00	55.41	O
ANISOU	2370	O	GLN	B	49	5795	7060	3155	1365	908	-1237 O
ATOM	2371	CB	GLN	B	49	86.704	-19.413	4.173	1.00	54.40	C
ANISOU	2371	CB	GLN	B	49	5403	7252	8533	1181	1015	-1440 C
ATOM	2372	CG	GLN	B	49	34.755	-10.326	3.054	1.09	52.37	C
ANISOU	2372	CG	GLN	B	49	7644	7552	9003	1564	1058	-1550 C
ATOM	2373	CD	GLN	B	49	85.386	-11.781	2.023	1.00	72.57	C
ANISOU	2373	CD	GLN	B	49	8370	8893	10304	1141	1185	-1583 C
ATOM	2374	OE1	GLN	B	49	84.672	-12.543	1.373	1.00	30.48	O
ANISOU	2374	OE1	GLN	B	49	9512	9752	11305	1082	1235	-1778 O
ATOM	2375	NE2	GLN	B	49	86.717	-11.751	1.389	1.03	13.02	N
ANISOU	2375	NE2	GLN	B	49	3315	9060	10370	1266	1245	-1697 N
ATOM	2376	N	THR	B	50	85.825	-3.585	7.439	1.06	54.26	N
ANISOU	2376	N	THR	B	50	5719	6391	8005	1154	774	-1133 N
ATOM	2377	CA	THR	B	50	80.350	-6.283	3.813	1.00	55.78	C
ANISOU	2377	CA	THR	B	50	5792	7204	5199	1237	720	-1045 C
ATOM	2378	C	THR	B	50	86.515	-8.839	9.754	1.05	51.07	C
ANISOU	2378	C	THR	B	50	6536	7310	8937	1275	674	-950 C
ATOM	2379	O	THR	B	50	84.859	-9.807	9.554	1.00	71.07	O
ANISOU	2379	O	THR	B	50	7953	8518	10231	1316	715	-965 O
ATOM	2380	CB	THR	B	50	86.645	-6.780	8.732	1.00	54.60	C
ANISOU	2380	CB	THR	B	50	5539	7234	7974	1101	675	-1037 C
ATOM	2381	OG1	THR	B	50	85.402	-6.012	8.790	1.00	54.53	O
ANISOU	2381	OG1	THR	B	50	5624	7150	7937	943	633	-1610 O
ATOM	2382	CG2	THR	B	50	87.473	-6.321	7.559	1.00	53.98	O
ANISOU	2382	CG2	THR	B	50	5376	7250	7355	1002	735	-1125 C
ATOM	2383	N	SER	B	51	85.495	-8.203	10.512	1.00	58.95	N
ANISOU	2383	N	SER	B	51	6244	7543	8612	1251	857	-863 N
ATOM	2384	CA	SER	B	51	84.615	-8.626	12.007	1.00	54.55	C
ANISOU	2384	CA	SER	B	51	5787	6359	8079	1265	559	-771 C
ATOM	2385	C	SER	B	51	83.835	-7.393	12.587	1.00	53.82	C
ANISOU	2385	C	SER	B	51	5597	5523	7932	1102	443	-733 C
ATOM	2386	O	SER	B	51	84.287	-6.270	12.294	1.00	53.90	O
ANISOU	2386	O	SER	B	51	5521	6972	7835	1057	435	-774 O

TABLE 3-continued

ATOM	2387	CB	SER	B	51	85.454	-9.237	13.112	1.0	52.30	C
ANISOU	2387	CB	SER	B	51	5519	6700	7879	1442	528	-674 C
ATOM	2388	OG	SER	B	51	88.342	-8.250	13.531	1.00	49.27	O
ANISOU	2388	OG	SER	B	51	4901	6471	7347	1410	466	-655 O
ATOM	2389	N	GLU	B	52	82.849	-7.812	13.401	1.60	52.14	N
ANISOU	2389	N	GLU	B	52	5536	6453	7737	1073	478	-572 N
ATOM	2390	CA	GLU	B	52	82.155	-6.512	14.127	1.00	54.79	C
ANISOU	2390	CA	GLU	B	52	5925	6354	8026	942	433	-634 C
ATOM	2391	C	GLU	B	52	83.134	-5.722	15.029	1.00	53.65	C
ANISOU	2391	C	GLU	B	52	5655	5907	7312	948	380	-598 C
ATOM	2392	O	GLU	B	52	83.062	-4.509	15.139	1.00	49.16	O
ANISOU	2392	O	GLU	B	52	5053	6423	7195	622	385	-618 O
ATOM	2393	CB	GLU	B	52	83.995	-7.042	14.977	1.00	52.99	C
ANISOU	2393	CB	GLU	B	52	5819	6484	7829	932	429	-564 C
ATCOM	2394	CG	GLU	B	52	73.843	-7.674	14.206	1.00	52.53	C
ANISOU	2394	CG	GLU	B	52	5883	6263	7345	879	478	-606 C
ATOM	2395	CD	GLU	B	52	78.035	-7.953	15.083	1.00	63.01	C
ANISOU	2395	CD	GLU	B	52	7295	7460	9164	533	477	-542 C
ATOM	2396	OE1	GLU	B	52	78.597	-7.648	16.300	1.30	56.29	O
ANISOU	2396	OE1	GLU	B	52	7705	7916	9558	646	445	-451 O
ATOM	2397	OE2	GLU	B	52	77.609	-8.479	14.570	1.00	70.17	O
ANISOU	2397	OE2	GLU	B	52	8283	8238	10140	771	513	-576 O
ATOM	2398	N	ALA	B	53	84.972	-5.429	15.644	1.00	50.11	N
ANISOU	2398	N	ALA	B	53	6422	7791	8625	1097	355	-549 N
ATOM	2399	CA	ALA	B	53	85.107	-5.805	16.471	1.00	53.79	C
ANISOU	2399	CA	ALA	B	53	5488	7205	7743	1108	296	-525 C
ATOM	2400	C	ALA	B	53	65.929	-4.791	15.579	1.00	54.13	C
ANISOU	2400	C	ALA	B	53	5406	7411	7758	1014	309	-515 C
ATOM	2401	O	ALA	B	53	86.311	-5.748	16.202	1.00	55.84	O
ANISOU	2401	O	ALA	B	53	5570	7898	5023	905	270	-829 O
ATOM	2402	CB	ALA	B	53	65.984	-8.891	17.080	1.00	58.50	C
ANISOU	2402	CB	ALA	B	53	5774	7595	5097	1315	269	-446 C
ATOM	2403	N	THR	B	54	36.21	5	-5.097	14.412	1.33	54.66 N
ANISOU	2403	N	THR	B	54	5461	7473	7891	1042	368	-585 N
ATOM	2404	CA	THR	B	54	35.969	-4.197	13.539	1.00	55.27	C
ANISOU	2404	CA	THR	B	54	5511	7757	7997	952	399	-772 C
ATOM	2405	C	THR	B	54	85.254	-2.916	13.138	1.00	52.79	C
ANISOU	2405	C	THR	B	54	5171	7324	7553	753	421	-810 C
ATOM	2406	O	THR	B	54	85.610	-1.621	12.971	1.90	47.05	O
ANISOU	2406	O	THR	B	54	4379	6732	6790	507	454	-854 O

TABLE 3-continued

ATOM	2407	CB	THR	B	54	87.330	-4.918	12.225	1.00	58.49	C
ANISOU	2407	CB	THR	B	54	5328	8029	8355	1030	470	-337 C
ATOM	2408	OG1	THR	B	54	83.041	-6.108	12.505	1.60	59.32	O
ANISOU	2408	OG1	THR	B	54	5950	8203	8577	1234	471	-802 O
ATOM	2409	CG2	THR	B	54	88.145	-4.014	11.271	1.00	52.36	C
ANISOU	2409	CG2	THR	B	54	4944	7401	7548	933	516	-924 C
ATOM	2410	N	VAL	B	55	84.943	-3.098	12.678	1.00	52.25	N
ANISOU	2410	N	VAL	B	55	5245	7379	7530	718	435	-793 N
ATOM	2411	CA	VAL	B	55	34.054	-1.988	12.520	1.00	47.95	C
ANISOU	2411	CA	VAL	B	55	4769	6468	8565	564	455	-801 C
ATOM	2412	C	VAL	B	55	80.974	-1.056	15.598	1.00	43.22	C
ANISOU	2412	C	VAL	B	55	4159	5941	5320	431	424	-771 C
ATOM	2413	O	VAL	B	55	84.057	0.149	13.523	1.00	50.54	O
ANISOU	2413	O	VAL	B	55	5585	5933	7222	354	452	-500 O
ATOM	2414	CB	VAL	B	55	82.885	-2.458	12.397	1.00	48.78	C
ANISOU	2414	CB	VAL	B	55	5003	8417	7115	559	466	-791 C
ATOM	2415	CG1	VAL	B	55	81.719	-1.304	11.994	1.00	43.71	C
ANISOU	2415	CG1	VAL	B	55	4419	5738	8451	424	479	-781 C
ATOM	2416	CG2	VAL	B	55	62.861	-3.237	10.810	1.00	47.49	C
ANISOU	2416	CG2	VAL	B	55	4548	6221	5974	603	505	-853 C
ATOM	2417	N	VAL	B	56	83.374	-1.629	14.899	1.00	44.57	N
ANISOU	2417	N	VAL	B	56	4554	6115	6504	549	374	-710 N
ATOM	2418	CA	VAL	B	56	63.669	-0.795	16.081	1.05	45.22	C
ANISOU	2418	CA	VAL	B	56	4560	6355	6648	462	348	-682 C
ATOM	2419	C	VAL	B	56	84.974	-0.022	16.254	1.00	49.83	C
ANISOU	2419	C	VAL	B	56	4561	7018	7037	401	339	-733 C
ATOM	2420	O	VAL	B	56	84.937	1.219	15.335	1.00	48.30	O
ANISOU	2420	O	VAL	B	56	515	4752	6925	6865	252	370 -774 O
ATOM	2421	CB	VAL	B	56	83.458	-1.636	17.347	1.00	47.25	C
ANISOU	2421	CB	VAL	B	56	4721	6452	5770	557	295	-602 C
ATOM	2422	CG1	VAL	B	56	83.569	-0.730	18.539	1.00	52.17	C
ANISOU	2422	CG1	VAL	B	56	5327	7184	7315	461	268	-593 C
ATOM	2423	CG2	VAL	B	56	82.148	-2.449	17.316	1.00	56.48	C
ANISOU	2423	CG2	VAL	B	56	5264	6570	7246	500	314	-552 CN
ATOM	2424	N	ARG	B	57	85.095	-0.765	16.273	1.00	50.73	N
ANISOU	2424	N	ARG	B	57	4075	7257	7144	517	306	-754 N
ATOM	2425	CA	ARG	B	57	37.421	-3.123	15.415	1.00	51.13	C
ANISOU	2425	CA	ARG	B	57	4763	7534	7131	460	292	-739 C
ATOM	2426	C	ARG	B	57	87.628	1.061	15.443	1.00	51.50	C
ANISOU	2426	C	ARG	B	57	4796	7605	7167	297	358	-874 C

TABLE 3-continued

ATOM	2427	O	ARG	B	57	38.027	2.141	15.345	1.00	50.35	O	
ANISOU	2427	O	ARG	B	57	4607	7562	6962	148	380	-920	
ATOM	2428	CB	ARG	B	57	88.521	-1.148	16.315	1.00	58.69	C	
ANISOU	2428	CB	ARG	B	57	5584	8616	8100	530	250	-776	
ATOM	2429	CG	ARG	B	57	89.793	-0.641	16.941	1.00	50.75	C	
ANISOU	2429	CG	ARG	B	57	5557	9144	8230	583	213	-807	
ATOM	2430	CD	ARG	B	57	90.928	-1.032	16.045	1.00	69.98	C	
ANISOU	2430	CD	ARG	B	57	5673	10441	9476	673	242	-853	
ATOM	2431	NE	ARG	B	57	91.314	-2.418	16.216	1.00	74.27	N	
ANISOU	2431	NE	ARG	B	57	7153	10891	10060	915	205	-782	
ATOM	2432	CZ	ARG	B	57	92.408	-2.931	15.664	1.00	82.84	C	
ANISOU	2432	CZ	ARG	B	57	6097	12210	11169	2037	223	-606	
ATOM	2433	NH1	ARG	B	57	93.189	-2.171	14.907	1.00	83.98	N	
ANISOU	2433	NH1	ARG	B	57	8116	12496	11295	922	276	-904	
ATOM	2434	NH2	ARG	B	57	92.718	-4.201	15.950	1.00	83.20	N	
ANISOU	2434	NH2	ARG	B	57	8114	12239	11259	1276	202	-731	
ATOM	2435	N	LEU	B	58	37.255	0.903	14.174	1.00	51.84	N	
ANISOU	2435	N	LEU	B	58	4897	7538	7261	311	428	-895	
ATOM	2436	CA	LEU	B	58	87.275	2.029	19.227	1.00	49.49	C	
ANISOU	2436	CA	LEU	B	58	4520	7236	5947	163	507	-952	
ATOM	2437	C	LEU	B	58	96.354	3.287	13.555	1.00	50.32	C	
ANISOU	2437	C	LEU	B	58	4845	7242	7082	8	543	-942	
ATOM	2438	O	LEU	B	58	85.732	4.491	13.443	1.00	48.03	O	
ANISOU	2438	O	LEU	B	58	4547	7013	6709	-145	602	-991	
ATOM	2439	CB	LEU	B	58	85.867	1.552	11.834	1.00	45.00	C	
ANISOU	2439	CB	LEU	B	58	4301	6753	8612	222	552	-960	
ATOM	2440	CG	LEU	B	58	85.815	2.647	10.922	1.00	54.16	C	
ANISOU	2440	CG	LEU	B	58	5308	7714	7555	82	632	-998	
ATOM	2441	CD1	LEU	B	58	38.205	0.251	11.019	1.00	55.44	C	
ANISOU	2441	CD1	LEU	B	58	5451	8105	7799	0	651	-1051	
ATOM	2442	CD2	LEU	B	58	86.475	2.144	9.512	1.00	51.92	C	
ANISOU	2442	CD2	LEU	B	58	5579	7359	7291	133	673	-1011	
ATOM	2443	N	CYS	B	59	85.111	3.006	13.933	1.00	45.37	N	
ANISOU	2443	N	CYS	B	59	4352	8421	5401	48	522	-961	
ATOM	2444	CA	CYS	B	59	84.194	4	048	14.379	1.00	43.80	
ANISOU	2444	CA	CYS	B	59	4261	5175	5244	-52	558	-363	
ATOM	2445	C	CYS	B	59	84.751	4.846	15.539	1.00	46.39	C	
ANISOU	2445	C	CYS	B	59	4533	6590	5498	-171	554	-895	
ATOM	2446	O	CYS	B	59	34.599	6.099	15.531	1.00	51.43	O	
ANISOU	8	2446	O	CYS	B	59	5230	7211	7119	-314	626	-930

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TABLE 3-continued

ATOM	2447	CB	CYS	B	59	82.845	3.394	14.724	1.00	43.94	C	
ANISOU	2447	CB	CYS	B	59	4372	5026	6298	18	527	-798	C
ATOM	2448	SG	CYS	B	59	82.045	2.759	13.210	1.00	44.22	S	
ANISOU	2448	SG	CYS	B	59	4465	5947	6390	88	546	-777	S
ATOM	2449	N	ARG	B	60	85.301	4.144	16.521	1.00	45.81	N	
ANISOU	2449	N	ARG	B	60	4511	5756	6520	-106	475	-955	N
ATOM	2450	CA	ARG	B	60	88.023	4.764	17.667	1.00	50.04	C	
ANISOU	2450	CA	ARG	B	60	5270	7733	7254	-203	4513	-923	C
ATOM	2451	C	ARG	B	60	87.240	5.680	37.173	1.50	55.93	C	
ANISOU	2451	C	ARG	B	60	5485	8218	7549	-348	500	-1022	C
ATOM	2452	O	ARG	B	60	87.421	5.763	17.627	1.00	54.39	O	
ANISOU	2452	O	ARG	B	60	5309	8055	7311	-519	548	-1034	O
ATOM	2453	CB	ARG	B	60	66.541	3.863	15.548	1.00	52.32	C	
ANISOU	2453	CB	ARG	B	60	5050	7730	7101	-79	349	-885	C
ATOM	2454	CG	ARG	B	60	85.863	3.474	19.881	1.00	54.26	C	
ANISOU	2454	CG	ARG	B	60	5089	7943	71305	-84	302	-831	C
ATOM	2455	CD	ARG	B	60	64.649	2.534	19.796	1.00	56.48	C	
ANISOU	2455	CD	ARG	B	60	5775	8024	7651	55	306	-746	C
ATOM	2456	NE	ARG	B	60	84.710	1.561	20.782	1.00	50.91	N	
ANISOU	2456	NE	ARG	B	60	7083	9387	8953	185	228	-668	N
ATOM	2457	CZ	ARG	B	60	83.664	0.868	21.263	1.00	72.54	C	
ANISOU	2457	CZ	ARG	B	60	7011	9053	3897	250	218	-585	C
ATOM	2458	NH1	ARG	B	60	82.407	1.131	20.887	1.00	72.15	N	
ANISOU	2458	NH1	ARG	B	60	7985	9715	9714	221	280	-579	N
ATOM	2459	NH2	ARG	B	60	83.882	-0.090	22.143	1.00	73.88	N	
ANISOU	2459	NH2	ARG	B	60	8067	10174	9829	383	142	-510	N
ATOM	2460	N	ASP	B	61	58.004	5.102	15.252	1.00	54.88	N	
ANISOU	2460	N	ASP	B	61	5241	8174	7438	-270	4618	-1040	C
ATOM	2461	CA	ASP	B	61	89.145	5.885	15.700	1.00	56.50	C	
ANISOU	2461	CA	ASP	B	61	5360	8539	7230	-409	557	-1128	C
ATOM	2462	C	ASP	B	61	88.677	7.352	14.969	1.00	5786	C	
ANISOU	2462	C	ASP	B	61	5635	8554	7793	-550	579	-1159	C
ATOM	2463	O	ASP	B	61	89.378	8.137	15.011	1.00	63.39	O	
ANISOU	2463	O	ASP	B	61	5294	9842	8449	-741	741	-1236	O
ATOM	2464	CB	ASP	B	61	90.005	3.010	14.787	1.00	69.44	C	
ANISOU	2464	CB	ASP	B	61	5578	3997	3011	-286	547	-1107	C
ATOM	2465	CG	ASP	B	61	90.885	3.408	15.554	1.00	63.84	C	
ANISOU	2465	CG	ASP	B	61	5737	9459	8350	-124	408	-1105	C
ATOM	2466	OD1	ASP	B	61	90.512	3.850	16.801	1.00	63.50	O	
ANISOU	2466	OD1	ASP	B	61	5938	9731	5459	-131	350	-1080	O

TABLE 3-continued

ATOM	2467	OD2	ASP	B	61	91.255	2.975	14.920	1.90	38.21	O
ANISOU	2467	OD2	ASP	B	61	5176	9084	7878	19	429	-1099 O
ATOM	2468	N	MET	B	62	87.500	7.146	14.328	1.00	58.23	N
ANISOU	2468	N	MET	B	62	5837	3694	7835	-515	715	-1007 N
ATOM	2469	CA	MET	B	62	85.951	8.358	13.738	1.00	54.05	C
ANISOU	2469	CA	MET	B	62	5443	7724	7373	-541	833	-1100 C
ATOM	2470	C	MET	B	62	86.341	9.383	14.7251	1.00	58.91	C
ANISOU	2470	C	MET	B	62	5167	8245	7989	-761	373	-1107 C
ATOM	2471	O	MET	B	62	85.848	13.457	14.339	1.00	57.96	O
ANISOU	2471	O	MET	B	62	5170	7990	1862	-854	950	-1101 O
ATOM	2472	CB	MET	B	62	85.303	5.015	12.657	1.00	55.82	C
ANISOU	2472	CB	MET	B	62	5772	7786	7651	-583	846	-1925 C
ATOM	2473	CG	MET	B	62	85.421	7.627	11.251	1.00	54.37	C
ANISOU	2473	CG	MET	B	62	5538	7551	7471	-490	573	-1037 C
ATOM	2474	SD	MET	B	62	67.785	8.740	13.704	3.00	7455	S
ANISOU	2474	SD	MET	B	62	8033	10344	9987	-674	987	-1123 S
ATOM	2475	CE	MET	B	62	86.925	10.2197	10.336	1.00	88.48	C
ANISOU	2475	CE	MET	B	62	7439	9364	9105	-609	1115	-1033 C
ATOM	2476	N	GLY	B	63	86.355	9.083	16.0213	1.00	32.13	N
ANISOU	2476	N	GLY	B	63	5271	7451	7073	-755	800	-1118 N
ATOM	2477	CA	GLY	B	63	85.745	9.974	17.010	1.30	54.53	C
ANISOU	2477	CA	GLY	B	63	5702	7677	7667	-864	848	-1134 C
ATOM	2478	C	GLY	B	63	84.388	9.570	17.598	1.00	54.73	C
ANISOU	2478	C	GLY	B	63	5832	7542	7421	-755	318	-1002 C
ATOM	2479	O	GLY	B	63	83.856	10.275	18.479	1.00	53.35	O
ANISOU	2479	O	GLY	B	63	5752	7293	7225	-838	365	-1069 O
ATOM	2480	N	TYR	B	64	33.845	8.423	17.178	1.00	50.03	N
ANISOU	2480	N	TYR	B	64	5234	5912	6884	-584	747	-974 N
ATOM	2481	CA	TYR	B	64	32.447	8.312	17.518	1.00	47.41	C
ANISOU	2481	CA	TYR	B	64	5008	5425	6602	-487	733	-893 C
ATOM	2482	C	TYR	B	64	82.418	7.017	18.6339	1.00	47.29	C
ANISOU	2482	C	TYR	B	64	4947	5463	6552	-406	637	-865 C
ATOM	2483	O	TYR	B	64	83.429	6.456	19.009	1.00	47.53	O
ANISOU	2483	O	TYR	B	64	4877	6575	6546	-384	566	-889 O
ATOM	2484	CB	TYR	B	64	61.681	7.511	16.256	1.00	43.31	C
ANISOU	2484	CB	TYR	B	64	4518	5790	6349	-377	735	-829 C
ATOM	2485	CG	TYR	B	64	81.583	3.535	15.230	1.00	46.41	C
ANISOU	2485	CG	TYR	B	64	4972	6109	6554	-456	837	-836 C
ATOM	2486	CD1	TYR	B	64	80.655	9.663	15.403	1.00	46.12	C
ANISOU	2486	CD1	TYR	B	64	5056	5928	6538	-504	924	-806 C

TABLE 3-continued

ATOM	2487	CD2	TYR	B	64	82.481	3.730	14.353	1.00	49.23	C
ANISOU	2487	CD2	TYR	B	64	5254	6533	6890	-453	851	-869 C
ATOM	2488	CE1	TYR	B	64	80.587	10.713	14.513	1.00	47.63	C
ANISOU	2488	CE1	TYR	B	64	5316	6343	6737	-563	1026	-706 C
ATOM	2489	CE2	TYR	B	64	82.415	9.797	13.265	1.00	46.64	C
ANISOU	2489	CE2	TYR	B	64	5011	6144	6567	-559	964	-863 C
ATOM	2490	CZ	TYR	B	64	51.454	10.765	13.435	1.00	47.75	C
ANISOU	2490	CZ	TYR	B	64	5281	6134	6731	-592	1043	-820 C
ATOM	2491	OH	TYR	B	64	81.338	11.842	12.634	1.00	53.92	O
ANISOU	2491	OH	TYR	B	64	6149	6825	7612	-650	1154	-794 O
ATOM	2492	N	LYS	B	65	81.253	6.794	19.217	1.00	45.60	N
ANISOU	2492	N	LYS	B	65	4952	6250	6492	-358	538	-809 N
ATOM	2493	CA	LYS	B	65	81.173	5.876	20.341	1.00	47.73	C
ANISOU	2493	CA	LYS	B	65	5077	6450	6593	-287	550	-772 C
ATOM	2494	C	LYS	B	65	81.123	4.434	19.576	1.00	47.93	C
ANISOU	2494	C	LYS	B	65	5064	5471	5576	-127	487	-709 C
ATOM	2495	O	LYS	B	65	81.379	3.509	20.546	1.00	51.28	O
ANISOU	2495	O	LYS	B	65	5456	6958	7069	-46	417	-671 O
ATOM	2496	CB	LYS	B	65	79.924	6.167	21.131	1.00	53.02	C
ANISOU	2496	CB	LYS	B	65	5869	6995	7282	-303	505	-736 C
ATOM	2497	CG	LYS	B	65	50.224	5.784	22.532	1.30	55.55	C
ANISOU	2497	CG	LYS	B	65	6213	7391	7537	-411	515	-735 C
ATOM	2498	CD	LYS	B	65	80.195	8.238	22.458	1.00	61.44	C
ANISOU	2498	CD	LYS	B	65	7020	8055	8239	-564	730	-811 C
ATOM	2499	CE	LYS	B	65	79.898	8.358	23.844	1.00	51.35	C
ANISOU	2499	CE	LYS	B	65	7085	3076	3150	-663	772	-902 C
ATOM	2500	NZ	LYS	B	65	75.720	9.737	23.712	1.00	54.71	N
ANISOU	2500	NZ	LYS	B	65	6374	7035	7373	-634	899	-892 N
ATOM	2501	N	GLY	B	66	80.748	4.222	15.523	1.00	45.69	N
ANISOU	2501	N	GLY	B	66	4792	5102	6464	-31	510	-597 N
ATOM	2502	CA	GLY	B	66	80.475	2.853	18.153	1.00	40.00	C
ANISOU	2502	CA	GLY	B	66	4067	5340	5802	55	462	-650 C
ATOM	2503	C	GLY	B	66	79.835	4.000	16.793	1.00	43.09	C
ANISOU	2503	C	GLY	B	66	4422	5632	6251	57	503	-651 C
ATOM	2504	O	GLY	B	66	79.791	4.114	16.393	1.00	41.18	O
ANISOU	2504	O	GLY	B	66	4257	5377	6003	-28	553	-577 O
ATOM	2505	N	TYR	B	67	79.337	1	875	16.231	1.00	39.95 N
ANISOU	2505	N	TYR	B	67	4311	5170	5910	152	476	-522 N
ATOM	2506	CA	TYR	B	67	78.821	1.6137	14.925	1.00	40.02	C
ANISOU	2506	CA	TYR	B	67	4136	5116	5955	157	499	-632 C

TABLE 3-continued

ATOM	2507	C	TYR	B	67	77.517	2.557	14.715	1.00	39.45	C
ANISOU	2507	C	TYR	B	67	4135	4929	5908	103	541	-599 C
ATOM	2508	O	TYR	B	67	77.449	3.386	13.747	1.00	42.58	O
ANISOU	2508	O	TYR	B	67	4536	5345	6296	57	576	-607 O
ATOM	2509	CB	TYR	B	67	78.596	0.414	14.453	1.90	39.10	C
ANISOU	2509	CB	TYR	B	67	4024	4949	5882	2	52	487 -627 C
ATOM	2510	CG	TYR	B	67	77.992	0.334	19.065	1.00	35.81	C
ANISOU	2510	CG	TYR	B	67	3629	4499	5438	240	485	-647 C
ATOM	2511	CD1	TYR	B	67	78.722	0.840	11.974	1.00	3468	C
ANISOU	2511	CD1	TYR	B	67	3437	4408	5301	215	508	-591 C
ATOM	2512	CD2	TYR	B	67	76.6159	-0.011	12.855	1.00	39.00	C
ANISOU	2512	CD2	TYR	B	67	4086	4796	6934	239	481	-620 C
ATOM	2513	CE1	TYR	B	67	78.155	0.853	10.690	1.30	35.59	C
ANISOU	2513	CE1	TYR	B	67	3537	4513	5424	292	520	-705 C
ATOM	2514	CE2	TYR	B	67	76.0133	0.019	11.573	1.09	35.90	C
ANISOU	2514	CE2	TYR	B	67	3829	4024	5669	219	485	-639 C
ATOM	2515	CZ	TYR	B	67	76.876	0.426	10.476	1.00	35.72	C
ANISOU	2515	CZ	TYR	B	67	3660	4450	5479	205	502	-650 C
ATOM	2516	OH	TYR	B	67	79.335	0.492	9.200	1.08	61.81	O
ANISOU	2516	OH	TYR	B	67	3309	4599	5568	134	503	-694 O
ATOM	2517	N	SER	B	68	76.511	2.490	15.593	1.09	38.89	N
ANISOU	2517	N	SER	B	68	4112	4801	5864	110	541	-554 N
ATOM	2518	CA	SER	B	68	75.283	3.247	15.547	1.90	35.24	C
ANISOU	2518	CA	SER	B	68	3827	4385	5558	77	585	-517 C
ATOM	2519	C	SER	B	68	75.4351	4.164	15.511	1.00	34.10	C
ANISOU	2519	C	SER	B	68	3078	4121	5258	5	1552	-526 C
ATOM	2520	O	SER	B	68	74.839	5.465	14.743	1.00	35.95	O
ANISOU	2520	O	SER	B	68	3837	4312	5510	-2	590	-498 O
ATOM	2521	CB	SER	B	68	74.341	2.801	16.715	1.00	32.81	C
ANISOU	2521	CB	SER	B	68	3431	3834	5151	91	535	-476 C
ATOM	2522	OG	SER	B	68	73.948	1.443	15.442	1.00	36.41	O
ANISOU	2522	OG	SER	B	68	3831	4305	5647	147	543	-456 O
ATOM	2523	N	ASP	B	69	76.401	5.247	15.3137	1.09	39.63	N
ANISOU	2523	N	ASP	B	69	4273	4577	5909	-50	661	-563 N
ATOM	2524	CA	ASP	B	69	75.726	6.573	16.421	1.00	39.27	C
ANISOU	2524	CA	ASP	B	69	4261	4827	5823	-143	746	-589 C
ATOM	2525	C	ASP	B	69	77.335	7.159	15.091	1.00	43.08	C
ANISOU	2525	C	ASP	B	69	4723	5340	6304	-166	772	-607 C
ATOM	2526	O	ASP	B	69	76.854	8.146	14.483	1.00	97.99	O
ANISOU	2526	O	ASP	B	69	4105	4626	5672	-191	544	-573 O

TABLE 3-continued

ATOM	2527	CB	ASP	B	69	77.714	6.822	17.564	1.00	41.86	C
ANISOU	2527	CB	ASP	B	69	4555	5243	6597	-208	737	-644 C
ATOM	2528	CG	ASP	B	69	73.047	8.265	17.375	1.00	47.55	C
ANISOU	2528	CB	ASP	B	69	5334	5950	6782	-332	830	-691 C
ATOM	2529	OD1	ASP	B	69	77.225	9.163	17.612	1.00	43.22	O
ANISOU	2529	OD1	ASP	B	69	4488	4904	5890	-349	916	-663 O
ATOM	2530	OD2	ASP	B	69	79.141	3.492	15.358	1.00	44.23	O
ANISOU	2530	OD2	ASP	B	69	4375	5646	6303	-412	821	-758 O
ATOM	2531	N	PHE	B	70	78.398	5.469	14.647	1.00	31.87	N
ANISOU	2531	N	PHE	B	70	3733	45132	5355	-150	722	-549 N
ATOM	2532	CA	PHE	B	70	78.081	5.729	13.313	1.00	37.05	C
ANISOU	2532	CA	PHE	B	70	3858	4722	5498	-185	747	-567 C
ATOM	2533	C	PHE	B	70	77.905	6.732	12.185	1.09	35.51	C
ANISOU	2533	C	PHE	B	70	3710	4452	5332	-114	754	-609 C
ATOM	2534	O	PHE	B	70	71.894	7.651	11.351	1.00	39.73	O
ANISOU	2534	O	PHE	B	70	4279	4962	5844	-151	817	-589 O
ATOM	2535	CB	PHE	B	70	80.075	5.660	12.977	1.00	33.50	C
ANISOU	2535	CB	PHE	B	70	3981	5059	5701	-118	667	-715 C
ATOM	2536	CG	PHE	B	70	50.592	5.773	11.575	1.00	42.35	C
ANISOU	2536	CG	PHE	B	70	4408	5553	6132	-126	715	-737 C
ATOM	2537	CD1	PHE	B	70	80.083	4.975	10.572	1.00	44.85	C
ANISOU	2537	CD1	PHE	B	70	4733	5845	6466	-51	687	-722 C
ATOM	2538	CD2	PHE	B	70	31.518	5.657	11.292	1.00	51.01	C
ANISOU	2588	CD2	PHE	B	70	5475	6722	7186	-220	779	-783 C
ATOM	2539	CE1	PHE	B	70	80.552	5.139	9.263	1.00	51.07	C
ANISOU	2539	CE1	PHE	B	70	5503	6480	7218	-54	720	-745 C
ATOM	2540	CE2	PHE	B	70	82.128	6.776	9.995	1.00	61.30	C
ANISOU	2540	CE2	PHE	B	70	6761	8070	8462	-284	818	-832 C
ATOM	2541	CZ	PHE	B	70	81.580	5.999	8.954	1.00	55.55	C
ANISOU	2541	CZ	PHE	B	70	6060	7330	7754	-150	737	-730 C
ATOM	2542	N	ARG	B	71	77.045	5.704	12.129	1.00	34.14	N
ANISOU	2542	N	ARG	B	71	3531	4245	5196	-35	593	-580 N
ATOM	2543	CA	ARG	B	71	76.035	5.587	11.067	1.00	34.17	C
ANISOU	2543	CA	ARG	B	71	3555	422	5214	6	582	-534 C
ATOM	2544	C	ARG	B	71	75.045	6.754	11.110	1.00	83.97	C
ANISOU	2544	C	ARG	B	71	3590	4112	5235	-5	740	-434 C
ATOM	2545	O	ARG	B	71	74.598	7.023	10.068	1.00	35.45	O
ANISOU	2545	O	ARG	B	71	3799	4296	5376	10	760	-416 O
ATOM	2546	CB	ARG	B	71	75.222	4.271	11.206	1.00	32.54	C
ANISOU	2546	CB	ARG	B	71	3711	4353	5430	67	613	-529 C

TABLE 3-continued

ATOM	2547	CG	ARG	B	71	74.501	5.832	9.378	1.00	38.62	C
ANISOU	2547	CG	ARG	B	71	4093	4770	5810	91	582	-522 C
ATOM	2548	CD	ARG	B	71	73.902	2.037	10.071	1.00	37.33	C
ANISOU	2548	CD	ARG	B	71	3922	4581	5453	124	526	-538 C
ATOM	2549	NE	ARG	B	71	72.687	2.790	10.843	1.30	89.84	N
ANISOU	2549	NE	ARG	B	71	4250	4835	5354	124	534	-480 N
ATOM	2550	CZ	ARG	B	71	71.532	2.127	10.753	1.00	43.58	C
ANISOU	2550	CZ	ARG	B	71	4710	5285	6554	131	503	-466 C
ATOM	2551	NH1	ARG	B	71	71.445	1.085	9.920	1.00	39.93	N
ANISOU	2551	NH1	ARG	B	71	4231	4846	5093	128	441	-516 N
ATOM	2552	NH2	ARG	B	71	70.448	2.523	11.475	1.30	38.37	N
ANISOU	2552	NH2	ARG	B	71	4051	4582	5947	132	523	-410 N
ATOM	2553	N	MET	B	72	74.503	7.157	12.312	1.00	25.21	N
ANISOU	2553	N	MET	B	72	8784	4217	5895	-22	772	-432 N
ATOM	2554	CA	MET	B	72	73.824	8.851	12.527	1.00	37.45	C
ANISOU	2554	CA	MET	B	72	4123	4406	5609	-24	851	-390 C
ATOM	2555	C	MET	B	72	74.510	9.600	11.920	1.00	37.17	C
ANISOU	2555	C	MET	B	72	4143	4355	5625	-79	939	-387 C
ATOM	2556	O	MET	B	72	73.883	10.891	11.177	1.00	87.80	O
ANISOU	2556	O	MET	B	72	4267	4384	5710	-45	989	-312 O
ATOM	2557	CB	MET	B	72	73.608	8574	14.039	1.00	34.14	C
ANISOU	2557	CB	MET	B	72	3755	3922	5302	-52	888	-436 C
ATOM	2558	CG	MET	B	72	72.692	9.756	14.333	1.00	37.213	C
ANISOU	2558	CG	MET	B	72	4203	4219	5730	-43	987	-353 C
ATOM	2559	SD	MET	B	72	70.903	9.456	14.063	1.00	39.97	S
ANISOU	2559	SD	MET	B	72	4518	4526	6142	55	954	-254 S
ATOM	2560	CE	MET	B	72	70.608	10.439	12.573	1.00	41.53	C
ANISOU	2560	CE	MET	B	72	4749	4719	62130	117	1305	-170 C
ATOM	2561	N	ALA	B	73	75.798	9.775	12.205	1.00	37.23	N
ANISOU	2561	N	ALA	B	73	4147	4419	5613	-163	960	-452 N
ATOM	2562	CA	ALA	B	73	75.513	10.933	11.515	1.00	37.24	C
ANISOU	2562	CA	ALA	B	73	4192	4397	5560	-269	1057	-459 C
ATOM	2563	C	ALA	B	73	76.587	10.834	10.097	1.00	89.07	C
ANISOU	2563	C	ALA	B	73	4414	4671	5759	-202	1043	-429 C
ATOM	2564	O	ALA	B	73	16.437	11.864	9.399	1.00	41.64	O
ANISOU	2564	O	ALA	B	73	4615	4937	6070	-212	1128	-369 O
ATOM	2565	CB	ALA	B	73	77.905	11.0.99	12.247	1.00	33.57	C
ANISOU	2565	CB	ALA	B	73	4329	4638	5668	-354	1079	-570 C
ATOM	2566	N	LEU	B	74	76.759	9.505	9.601	1.30	37.40	N
ANISOU	2566	N	LEU	B	74	4123	4552	5535	-155	946	-455 N

TABLE 3-continued

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ATOM	2567	CA	LEU	B	74	76.394	9.381	3.188	1.00	39.66	C
ANISOU	2567	CA	LEU	B	74	4397	4897	5775	-129	929	-438 C
ATOM	2568	C	LEU	B	74	75.555	9.793	7.552	1.00	39.33	C
ANISOU	2568	C	LEU	B	74	4404	4802	5739	-57	929	-329 C
ATOM	2569	O	LEU	B	74	75.543	10.508	6.553	1.00	38.40	O
ANISOU	2569	O	LEU	B	74	4333	4683	5574	-54	977	-270 C
ATOM	2570	CB	LEU	B	74	77.246	7.943	7.874	1.00	37.57	C
ANISOU	2570	CB	LEU	B	74	4050	4723	5501	-91	836	-500 C
ATOM	2571	CG	LEU	B	74	77.169	7.6124	5.373	1.00	43.26	C
ANISOU	2571	CG	LEU	B	74	4767	5505	6165	-63	316	-488 C
ATOM	2572	CD1	LEU	B	74	78.235	3.417	5.573	1.00	45.21	C
ANISOU	2572	CD1	LEU	B	74	5034	5797	5347	-131	900	-802 C
ATOM	2573	CD2	LEU	B	74	77.322	6.152	6.129	1.00	42.33	C
ANISOU	2573	CD2	LEU	B	74	4719	5581	5184	-21	725	-557 C
ATOM	2574	N	ALA	B	75	74.431	9.337	8.189	1.00	39.65	N
ANISOU	2574	N	ALA	B	75	4429	4604	5384	4	380	-295 N
ATOM	2575	CA	ALA	B	75	73.125	9	773	7.585	1.00	39.16 C
ANISOU	2575	CA	ALA	B	75	4384	4715	5779	81	374	-188 C
ATOM	2576	C	ALA	B	75	72.876	11.284	7.569	1.00	36.33	C
ANISOU	2576	C	ALA	B	75	4371	4612	5580	92	988	-99 C
ATOM	2577	O	ALA	B	75	72.377	11.318	6.593	1.00	39.47	C
ANISOU	2577	O	ALA	B	75	4544	4154	5789	148	1002	-3 C
ATOM	2578	CB	ALA	B	75	72.013	9.039	8.342	1.00	37.01	C
ANISOU	2578	CB	ALA	B	75	4053	4428	5672	130	811	-180 C
ATOM	2579	N	VAL	B	76	73.245	11.981	8.639	1.00	37.71	N
ANISOU	2579	N	VAL	B	76	4347	4340	5640	36	1076	-130 N
ATOM	2580	CA	VAL	B	76	72.988	13.425	8.742	1.00	41.28	C
ANISOU	2580	CA	VAL	B	76	4908	4665	6112	42	1210	-56 C
ATOM	2581	C	VAL	B	76	73.873	14.073	7.576	1.00	46.17	C
ANISOU	2581	C	VAL	B	76	5584	5276	6602	-8	1273	-39 C
ATOM	2582	O	VAL	B	76	73.443	14.951	6.904	1.00	44.98	C
ANISOU	2582	O	VAL	B	76	5509	5088	6494	48	1341	75 C
ATOM	2583	CB	VAL	B	76	73.815	14.001	10.155	1.00	41.00	C
ANISOU	2583	CB	VAL	B	76	4929	4530	6120	-38	1300	-125 C
ATOM	2584	CG1	VAL	B	76	73.304	15.529	10.139	1.00	45.64	C
ANISOU	2584	CG1	VAL	B	76	5652	4967	6721	-58	1465	-71 C
ATOM	2585	CG	VAL	B	76	2	72.264	10.522	11.176	1.00	43.25 C
ANISOU	2585	CG2	VAL	B	76	5175	4788	6469	23	1252	-119 C
ATOM	2586	N	ASP	B	77	75.122	13.530	7.635	1.00	45.58	N
ANISOU	2586	N	ASP	B	77	5593	5426	6673	-109	1257	-145 N

TABLE 3-continued

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ATOM	2587	CA	ASP	B	77	75.055	14.101	5.599	1.00	52.16	C
ANISOU	2587	CA	ASP	B	77	6242	5166	7310	-172	1317	-143 C
ATOM	2588	C	ASP	B	77	75.517	13.957	5.161	1.00	49.49	C
ANISOU	2588	C	ASP	B	77	6006	5889	5908	-82	1274	-42 C
ATOM	2589	O	ASP	B	77	75.467	14.941	4.444	1.00	52.20	C
ANISOU	2589	O	ASP	B	77	5571	6301	7042	-72	1355	52 C
ATOM	2590	CB	ASP	B	77	77.358	13.393	6.769	1.00	54.12	C
ANISOU	2590	CB	ASP	B	77	5508	5525	7529	-270	1283	-275 C
ATOM	2591	CG	ASP	B	77	78.501	14.299	6.641	1.00	65.00	C
ANISOU	2591	CG	ASP	B	77	7939	7888	8871	-397	1404	-315 C
ATOM	2592	OD1	ASP	B	77	79.104	14.759	7.531	1.00	68.56	O
ANISOU	2592	OD1	ASP	B	77	8401	8302	9345	-504	1467	-392 O
ATOM	2593	OD2	ASP	B	77	78.745	14.559	5.356	1.00	63.95	O
ANISOU	2593	OD2	ASP	B	77	7839	7782	8677	-398	1442	-266 O
ATOM	2594	N	LEU	B	78	75.038	12.755	4.740	1.01	48.18	N
ANISOU	2594	N	LEU	B	78	5745	5808	5723	-21	1139	-59 N
ATOM	2595	CA	LEU	B	78	74.517	12.610	3.396	1.00	46.92	C
ANISOU	2595	CA	LEU	B	78	5584	5758	6487	53	1059	28 C
ATOM	2596	C	LEU	B	78	78.177	13.346	3139	1.00	49.87	C
ANISOU	2596	C	LEU	B	78	6000	6077	6570	170	1098	165 C
ATOM	2597	O	LEU	B	78	72.354	13.358	2.308	1.00	52.20	O
ANISOU	2597	O	LEU	B	78	6324	5425	7034	226	1091	280 O
ATOM	2598	CB	LEU	B	78	74.358	11.132	3.040	1.00	47.82	C
ANISOU	2598	CB	LEU	B	78	5592	6002	5577	72	952	-47 C
ATOM	2599	CG	LEU	B	78	75.644	10.305	2.996	1.00	49.92	C
ANISOU	2599	CG	LEU	B	78	5810	6345	6813	-6	936	-181 C
ATOM	2600	CD1	LEU	B	78	75.260	8.852	3.111	1.00	45.43	C
ANISOU	2600	CD1	LEU	B	78	5275	5950	6399	25	822	-258 C
ATOM	2601	CD2	LEU	B	78	76.244	10.500	1.515	1.00	51.59	C
ANISOU	2601	CD2	LEU	B	78	6039	5675	5950	-31	973	-157 C
ATOM	2602	N	SER	B	79	72.381	13.612	4.105	1.00	48.45	C
ANISOU	2602	N	SER	B	79	5834	5858	6847	218	1114	214 N
ATOM	2603	CA	SER	B	79	71.127	14.333	3.920	1.00	53.19	N
ANISOU	2603	CA	SER	B	79	6445	3361	7401	847	1132	869 C
ATOM	2604	C	SER	B	79	71.412	15.813	3.723	1.06	60.06	C
ANISOU	2604	C	SER	B	79	7500	7128	3301	356	1292	458 C
ATOM	2605	O	SER	B	79	73.589	15.501	3.165	1.00	71.15	O
ANISOU	2605	O	SER	B	79	8906	8476	9654	470	1315	522 O
ATOM	2606	CB	SER	B	79	73.125	14.685	5.054	1.00	52.09	C
ANISOU	2606	CB	SER	B	79	7139	1061	8251	402	3105	355 C

TABLE 3-continued

ATOM	2607	OG	SER	B	79	70.478	14.809	6.236	1	00	59.27	O
ANISOU	2607	OG	SER	B	79	7330	7018	8439	351	1227	324	O
ATOM	2608	N	GLN	B	80	72.509	16.270	4.092	1.00	62.40	N	
ANISOU	2608	N	GLN	B	80	7841	7311	8555	225	1400	383	N
ATOM	2609	CA	GLN	B	80	73.029	17.578	3.830	1.03	69.76	C	
ANISOU	2609	CA	GLN	B	80	8925	5093	9485	200	1574	461	C
ATOM	2610	C	GLN	B	80	73.595	17.957	2.484	1.30	65.08	C	
ANISOU	2610	C	GLN	B	80	8510	7605	8906	171	1506	510	C
ATOM	2611	O	GLN	B	80	73.776	17.093	1.605	1.00	72.54	O	
ANISOU	2611	O	GLN	B	80	9301	8721	9690	184	1489	504	O
ATOM	2612	CB	GLN	B	80	73.902	18.199	4.978	1.00	74.50	C	
ANISOU	2612	CB	GLN	B	80	9504	0581	10160	55	1595	338	C
ATOM	2613	CG	GLN	B	80	73.088	18.504	5.222	1.60	81.05	C	
ANISOU	2613	CG	GLN	B	80	10515	9353	11157	103	1734	336	C
ATOM	2614	CD	GLN	B	80	73.942	18.829	7	423	1.00	92.72	C
ANISOU	2614	CD	GLN	B	80	11961	10670	12599	-65	1625	190	C
ATOM	2615	OE1	GLN	B	80	73.739	19.858	8.078	1.00	104.45	O	
ANISOU	2615	OE1	GLN	B	80	13566	11981	14143	-72	1973	204	O
ATOM	2616	NE2	GLN	B	80	74.907	17.904	7.720	1.00	86.87	N	
ANISOU	2616	NE2	GLN	B	80	11503	10437	12206	-174	1743	48	N
ATOM	2617	N	ASP	B	92	52.904	10.863	-5.324	1.00	52.01	N	
ANISOU	2617	N	ASP	B	92	7323	6789	5651	912	1422	1097	N
ATOM	2618	CA	ASP	B	92	51.921	14.227	-5.313	1.00	63.42	C	
ANISOU	2618	CA	ASP	B	92	5763	9171	7102	925	1250	1007	C
ATOM	2619	C	ASP	B	92	51.962	13.397	-3980	1.00	66.46	C	
ANISOU	2619	C	ASP	B	92	9070	8517	7658	867	1283	980	C
ATOM	2620	O	ASP	B	92	10.952	12.176	-4.021	1.00	67.52	O	
ANISOU	2620	O	ASP	B	92	9153	8721	7767	860	1225	869	O
ATOM	2621	CB	ASP	B	92	56.502	14.096	-5.875	1.00	56.86	C	
ANISOU	2621	CB	ASP	B	92	8457	7926	6723	1014	1267	1120	C
ATOM	2622	CG	ASP	B	92	49.439	14.423	-4.813	1.00	65.27	C	
ANISOU	2622	CG	ASP	B	92	9013	8435	7355	1030	1193	1115	C
ATOM	2623	OD1	ASP	B	92	49.313	15.525	-4.511	1.33	64.95	O	
ANISOU	2623	OD1	ASP	B	92	90085	8308	7361	1043	1243	1198	O
ATOM	2624	OD2	ASP	B	92	48.760	13.540	-4.237	1.00	67.56	O	
ANISOU	2624	OD2	ASP	B	92	9249	8763	7657	1023	1099	1029	O
ATOM	2625	N	ILE	B	93	51.918	14.067	-2.819	1.05	61.50	N	
ANISOU	2625	N	ILE	B	93	8440	7782	7145	833	1282	987	N
ATOM	2626	CA	ILE	B	93	52.083	13.395	-3.521	3.00	61.50	C	
ANISOU	2626	CA	ILE	B	93	8375	7741	7249	775	1227	869	C

TABLE 3-continued

ATOM	2627	C	ILE	B	93	51.237	32.156	-1.289	1.00	53.64	C	
ANISOU	2627	C	ILE	B	93	7339	6821	6220	794	1119	903	C
ATOM	2628	O	ILE	B	93	51.306	11.143	-0.923	1.00	51.92	O	
ANISOU	2628	O	ILE	B	93	7072	5517	5938	749	1097	717	O
ATOM	2629	CB	ILE	B	93	51.909	14.325	-0.310	1.00	61.68	C	
ANISOU	2629	CB	ILE	B	93	8403	7853	7383	748	1229	908	C
ATOM	2630	CG1	ILE	B	93	53.013	15.374	-0.306	1.00	65.63	C	
ANISOU	2630	CG1	ILE	B	93	9056	8185	8075	700	1336	965	C
ATOM	2631	CG2	ILE	B	93	52.014	13.520	0.931	1.00	65.25	C	
ANISOU	2631	CG2	ILE	B	93	8915	3206	8050	594	1167	816	C
ATOM	2632	CD1	ILE	B	93	53.510	15.718	1.083	1.00	7067	C	
ANISOU	2632	CD1	ILE	B	93	9535	8894	8723	529	1339	922	C
ATOM	2633	N	CYS	B	94	49.921	12.204	-1.473	1.00	53.55	N	
ANISOU	2633	N	CYS	B	94	7344	6357	5145	857	1053	625	N
ATOM	2634	CA	CYS	B	94	49.125	11.004	-1.210	1.00	53.34	C	
ANISOU	2634	CA	CYS	B	94	7272	5399	6096	864	951	741	C
ATOM	2635	C	CYS	B	94	49.850	9.355	-2.031	1.00	54.96	C	
ANISOU	2635	C	CYS	B	94	7463	7190	6230	861	941	880	C
ATOM	2636	O	CYS	B	94	49.525	8.897	-1.615	1.00	47.33	O	
ANISOU	2636	O	CYS	B	94	6448	5243	5294	625	385	586	O
ATOM	2637	CB	CYS	B	94	47.654	11.231	-1.402	1.05	57.31	C	
ANISOU	2637	CB	CYS	B	94	7788	7452	6535	933	888	775	C
ATOM	2638	SG	CYS	B	94	47.143	12.712	-0.525	1.00	79.30	S	
ANISOU	2638	SG	CYS	B	94	10612	10146	91407	953	905	853	S
ATOM	2639	N	ASP	B	95	40.317	10.173	-3.349	1.00	50.78	N	
ANISOU	2639	N	ASP	B	95	5977	5714	5503	901	995	735	N
ATOM	2640	CA	ASP	B	95	50.105	9.116	-4.314	1.00	53.25	C	
ANISOU	2640	CA	ASP	B	95	7283	7123	5828	913	984	677	C
ATOM	2641	C	ASP	B	95	51.492	8.530	-4.125	1.00	49.40	C	
ANISOU	2641	C	ASP	B	95	5762	5604	5402	850	1038	616	C
ATOM	2642	O	ASP	B	95	51.577	7.325	-4.286	1.00	59.14	O	
ANISOU	2642	O	ASP	B	95	6824	8745	5481	338	998	523	O
ATOM	2643	CB	ASP	B	95	49.931	9.616	-5.742	1.00	62.50	C	
ANISOU	2643	CB	ASP	B	95	8512	8372	8863	976	1025	753	C
ATOM	2644	CG	ASP	B	95	43.597	10.315	-5.055	1.09	64.88	C	
ANISOU	2644	CG	ASP	B	95	8804	8651	7002	1045	976	827	C
ATOM	2645	OD1	ASP	B	95	48.676	13.493	-5.359	1.00	72.78	O	
ANISOU	2645	OD1	ASP	B	95	2599	9674	3081	1077	1041	936	O
ATOM	2646	OD2	ASP	B	95	47.508	9.705	-5.736	1.00	81.14	O	
ANISOU	2646	OD2	ASP	B	95	8344	8278	6609	1067	876	779	O

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TABLE 3-continued

ATOM	2647	N	VAL	B	96	52.452	9.295	-3.821	1.00	45.09	N
ANISOU	2647	N	VAL	B	96	5224	5982	4928	813	1127	565 N
ATOM	2648	CA	VAL	B	96	53.362	8.947	-3	576	1.00	46.20 C
ANISOU	2648	CA	VAL	B	96	6324	6953	5188	753	1163	512 C
ATOM	2649	C	VAL	B	96	53.911	8.089	-2.294	1.00	41.20	C
ANISOU	2649	C	VAL	B	96	5630	5413	4611	704	1113	513 C
ATOM	2650	O	VAL	B	96	54.547	7.011	-2.212	1.09	45.00	O
ANISOU	2650	O	VAL	B	96	5195	5040	5242	580	1098	433 O
ATOM	2651	CB	VAL	B	96	54.832	10.153	-3.521	1.50	47.15	C
ANISOU	2651	CB	VAL	B	96	6463	6136	8315	718	1293	622 C
ATOM	2652	CG1	VAL	B	96	55.208	9.764	-3.925	1.50	47.30	C
ANISOU	2652	CG1	VAL	B	96	5426	5119	5427	650	1340	634 C
ATOM	2653	CG2	VAL	B	96	54.998	10.798	-4.932	1.00	45.91	C
ANISOU	2653	CG2	VAL	B	96	5744	6417	5424	761	1379	780 C
ATOM	2654	N	SER	B	97	63.182	6.535	-1.296	1.30	37.85	N
ANISOU	2654	N	SER	B	97	5140	4869	4184	696	1066	534 N
ATOM	2655	CA	SER	B	97	53.139	7.755	-0.002	1.30	93.28	C
ANISOU	2655	CA	SER	B	97	6202	4945	4396	550	996	453 C
ATOM	2656	C	SER	B	97	52.545	6.379	-0.251	1.00	37.87	C
ANISOU	2656	C	SER	B	97	5130	4965	4292	670	913	374 C
ATOM	2657	O	SER	B	97	53.038	5.344	0.205	1.00	37.45	O
ANISOU	2657	O	SER	B	97	5099	4910	4296	535	882	293 O
ATOM	2658	CB	SER	B	97	52.225	8.467	0.952	1.00	35.18	C
ANISOU	2658	CB	SER	B	97	4316	4499	4050	651	967	437 C
ATOM	2659	OG	SER	B	97	52.714	5.769	1.204	1.00	46.10	O
ANISOU	2659	OG	SER	B	97	6225	5807	5483	634	1092	557 O
ATOM	2660	N	ALA	B	98	61.474	6.365	-1.033	1.00	39.57	N
ANISOU	2660	N	ALA	B	98	5389	5265	4417	726	574	997 N
ATOM	2661	CA	ALA	B	98	50.787	5.199	-1.370	1.00	41.74	C
ANISOU	2661	CA	ALA	B	98	5534	6596	4629	741	769	320 C
ATOM	2662	C	ALA	B	98	51.619	4.207	-2.237	1.00	41.21	C
ANISOU	2662	C	ALA	B	98	5562	5578	4517	743	814	261 C
ATOM	2663	O	ALA	B	98	51.610	2.992	-2.016	1.00	41.37	O
ANISOU	2663	O	ALA	B	98	5552	7609	4.557	725	757	171 O
ATOM	2664	CB	ALA	B	98	43.421	5.319	-2.008	1.00	43.48	C
ANISOU	2664	CB	ALA	B	98	5861	5888	4750	799	706	355 C
ATOM	2665	N	GLN	B	99	52.316	4.778	-3.232	1.00	44.06	N
ANISOU	2665	N	GLN	B	99	5956	5968	4817	765	900	307 N
ATOM	2666	CA	GLN	B	99	53.086	1.973	-4.231	1.00	40.06	C
ANISOU	2666	CA	GLN	B	99	5448	5522	4254	777	939	251 C

TABLE 3-continued

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ATOM	2667	C	GLN	B	99	54.354	3.326	-3.491	1.33	42.57	C
ANISOU	2667	C	GLN	B	99	5715	5782	4677	725	958	184 C
ATOM	2668	O	GLN	B	99	64.647	2.166	-3.666	1.00	40.05	O
ANISOU	2668	O	GLN	B	99	5376	5491	4351	724	931	O
ATOM	2669	CB	GLN	B	99	53.573	4844	-5.398	1.00	44.39	C
ANISOU	2669	CB	GLN	B	99	6039	6113	4703	312	1031	329 C
ATOM	2670	CG	GLN	B	99	54.311	4.033	-6.445	1.00	45.63	C
ANISOU	2670	CG	GLN	B	99	6200	6349	4790	830	1068	269 C
ATOM	2671	CD	GLN	B	99	53.465	2.920	-7.023	1.93	51.40	C
ANISOU	2671	CD	GLN	B	99	6936	7159	5434	855	880	189 C
ATOM	2672	OE1	GLN	B	99	52.420	3.192	-7.645	1.00	50.23	O
ANISOU	2672	OE1	GLN	B	99	6824	7075	5185	910	939	225 O
ATOM	2673	NE2	GLN	B	99	53.937	1.675	-6.889	1.00	52.16	N
ANISOU	2673	NE2	GLN	B	99	6998	7256	5566	847	951	78 N
ATOM	2674	N	SER	B	100	54.937	4.112	-2.544	1.50	42.03	N
ANISOU	2674	N	SER	B	100	5630	5636	4706	684	1006	227 N
ATOM	2675	CA	SER	B	100	56.010	3.624	-1.369	1.00	45.37	C
ANISOU	2675	CA	SER	B	100	6063	6067	5297	635	1021	173 C
ATOM	2576	C	SER	B	100	56.560	2.412	-0.994	1.00	42.44	C
ANISOU	2676	C	SER	B	100	5594	5513	4919	619	919	66 C
ATOM	2677	O	SER	B	100	55.299	1.435	-0.871	1.00	39	10 O
ANISOU	2677	O	SER	B	100	5135	5135	4535	605	912	12 O
ATOM	2678	CB	SER	B	100	56.502	4.513	-1.097	1.00	51.13	C
ANISOU	2678	CB	SER	B	100	6721	6655	6051	594	1081	239 C
ATOM	2679	OG	SER	B	100	57.730	4.446	-0.406	1.00	60.02	O
ANISOU	2679	OG	SER	B	100	7792	7736	7275	543	1102	192 O
ATOM	2680	N	ALA	B	101	54.332	2.418	-0.474	1.00	43.43	N
ANISOU	2680	N	ALA	B	101	5725	5732	5041	629	841	95 N
ATOM	2681	CA	ALA	B	101	53.854	1.255	0.310	1.00	33.76	C
ANISOU	2681	CA	ALA	B	101	5106	5121	4501	6011	747	22 C
ATOM	2682	C	ALA	B	101	53.426	0.071	-0.530	1.00	38.90	C
ANISOU	2682	C	ALA	B	101	5133	5206	4442	631	896	-52 C
ATOM	2683	O	ALA	B	101	53.647	-1.090	-0.201	1.00	40.06	O
ANISOU	2683	O	ALA	B	101	5253	5336	4832	615	651	-131 O
ATOM	2684	CB	ALA	B	101	52.720	1.757	1.250	1.00	37.53	C
ANISOU	2684	CB	ALA	B	101	4952	4938	4370	590	689	59 C
ATOM	2685	N	VAL	B	102	52.853	0.352	-1.686	1.00	39.34	N
ANISOU	2685	N	VAL	B	102	5228	5337	4382	577	705	-28 N
ATOM	2686	CA	VAL	B	102	52.557	-0.702	-2.620	1.00	38.72	C
ANISOU	2686	CA	VAL	B	102	5161	5327	4224	705	666	-102 C

TABLE 3-continued

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ATOM	2687	C	VAL	B	102	53.860	-1.405	-3.032	1.00	42.14	C
ANISOU	2687	C	VAL	B	102	5579	5759	4674	705	718	-186 C
ATOM	2688	O	VAL	B	102	53.923	-2.644	-3.057	1.00	42.72	O
ANISOU	2688	O	VAL	B	102	5638	5832	4751	702	670	-258 O
ATOM	2689	CB	VAL	B	102	51.851	-0.088	-3.888	1.00	39.55	C
ANISOU	2689	CB	VAL	B	102	5314	5522	4195	753	078	-53 C
ATOM	2690	CG1	VAL	B	102	51.922	-1.051	-3.036	1.00	33.32	C
ANISOU	2690	CG1	VAL	B	102	5174	5445	3941	791	667	-131 C
ATOM	2691	CG2	VAL	B	102	50.398	0.283	-3.451	1.00	41.49	C
ANISOU	2691	CG2	VAL	B	102	6582	5778	4422	764	607	-14 C
ATOM	2692	N	ASP	B	103	54.910	-0.532	-0.320	1.00	38.49	N
ANISOU	2692	N	ASP	B	103	3117	3292	4216	707	817	-119 N
ATOM	2693	CA	ASP	B	103	56.203	-1.218	-3.717	1.00	44.15	C
ANISOU	2693	CA	ASP	B	103	5811	5015	4949	711	877	-175 C
ATOM	2694	C	ASP	B	103	55.791	-2.140	-2.674	1.00	41.55	C
ANISOU	2694	C	ASP	B	103	5431	5614	4741	675	839	-245 C
ATOM	2695	O	ASP	B	103	37.268	-3.265	-2.985	1.00	38.72	O
ANISOU	2695	O	ASP	B	103	6058	5255	4386	591	828	-334 O
ATOM	2696	CB	ASP	B	103	57.204	-0.109	-4.032	1.00	44.63	C
ANISOU	2696	CB	ASP	B	103	5874	6079	5006	708	993	-101 C
ATOM	2697	CG	ASP	B	103	56.825	0.607	-5.384	1.00	49.17	C
ANISOU	2697	CG	ASP	B	103	6504	6737	5443	753	1042	-40 C
ATOM	2698	OD1	ASP	B	103	54.555	0.104	-5.147	1.00	45	36 O
ANISOU	2698	OD1	ASP	B	103	6053	6323	4850	792	991	-72 O
ATOM	2699	OD2	ASP	B	103	57.364	1.677	-5.661	1.00	51.38	O
ANISOU	2699	OD2	ASP	B	103	5795	7015	5713	749	1132	42 O
ATOM	2700	N	SER	B	104	55.752	-1.691	-1.428	1.00	40.53	N
ANISOU	2700	N	SER	B	104	5315	5451	4747	634	326	-267 N
ATOM	2701	CA	SER	B	104	57.258	-2.487	-0.316	1.00	42.41	C
ANISOU	2701	CA	SER	B	104	5468	5583	5062	600	779	-253 C
ATOM	2702	C	SER	B	104	56.428	-3.733	-0.103	1.00	44.05	C
ANISOU	2702	C	SER	B	104	5081	5785	5272	502	579	-331 C
ATOM	2703	O	SER	B	104	57.000	-4.755	0.223	1.00	42.74	O
ANISOU	2703	O	SER	B	104	6488	5586	5167	598	655	-398 O
ATOM	2704	CB	SER	B	104	57.271	-1.718	1.002	1.00	47.10	C
ANISOU	2704	CB	SER	B	104	6039	6108	5749	554	771	-204 C
ATOM	2705	OG	SER	B	104	57.993	-3.514	3.329	1.00	54.00	O
ANISOU	2705	OG	SER	B	104	5912	6979	6527	545	362	-137 O
ATOM	2706	N	LEU	B	105	55.094	-3.040	-0.196	1.00	35.7	N
ANISOU	2706	N	LEU	B	105	5150	5260	4567	535	513	-313 N

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ATOM	2707	CA	LEU	B	105	54.294	-4.368	-0.161	1.30	27.08	C
ANISOU	2707	CA	LEU	B	105	4907	5005	4405	504	527	-385 C
ATOM	2708	C	LEU	B	105	54.085	-5.850	-1.248	1.00	40.18	C
ANISOU	2708	C	LEU	B	105	5238	5363	4667	539	533	-472 C
ATOM	2709	O	LEU	B	105	54.843	-7.653	-1.026	1.00	38.88	O
ANISOU	2709	O	LEU	B	105	5061	5163	4546	533	487	-551 O
ATOM	2710	CB	LEU	B	105	52.848	-4.522	-0.367	1.00	35.21	C
ANISOU	2710	CB	LEU	B	105	4520	4735	4022	608	473	-352 C
ATOM	2711	CG	LEU	B	105	52.142	-3.870	0.832	1.00	33.43	C
ANISOU	2711	CG	LEU	B	105	4379	4466	3855	574	440	-289 C
ATOM	2712	CD1	LEU	B	105	50.756	-3.371	0.367	1.00	34.24	C
ANISOU	2712	CD1	LEU	B	105	4506	4631	3871	504	401	-251 C
ATOM	2713	CD2	LEU	B	105	52.063	-4.775	2.053	1.00	35.86	C
ANISOU	2713	CD2	LEU	B	105	4656	4706	4263	529	375	-328 C
ATOM	2714	N	GLN	B	106	54.831	-5.346	-2.461	1.00	38.14	N
ANISOU	2714	N	GLN	B	106	5008	5180	4305	679	592	-456 N
ATOM	2715	CA	GLN	B	106	55.232	-6.236	-3.533	1.00	41.71	C
ANISOU	2715	CA	GLN	B	106	5475	5579	4694	716	695	-542 C
ATOM	2716	C	GLN	B	106	56.585	-6.846	-3.310	1.00	41.52	C
ANISOU	2716	C	GLN	B	106	5416	5611	4748	719	649	-596 C
ATOM	2717	O	GLN	B	106	55.757	-6.061	-3.424	1.00	45.58	O
ANISOU	2717	O	GLN	B	106	5940	6119	5298	730	610	-691 O
ATOM	2718	CB	GLN	B	106	55.107	-5.594	-4.365	1.00	44.10	C
ANISOU	2718	CB	GLN	B	106	5817	6080	4860	759	663	-507 C
ATOM	2719	CG	GLN	B	106	52.521	-5.325	-5.119	1.00	44.44	C
ANISOU	2719	CG	GLN	B	106	6590	6159	4827	764	559	-455 C
ATOM	2720	CD	GLN	B	106	53.296	-4.402	-6.243	1.00	46.16	C
ANISOU	2720	CD	GLN	B	106	6145	6482	4912	306	534	-423 C
ATOM	2721	OE1	GLN	B	106	54.161	-3.729	-6.795	1.00	51.25	O
ANISOU	2721	OE1	GLN	B	106	6814	7163	5323	827	729	-879 O
ATOM	2722	NE2	GLN	B	106	52.044	-4.375	-6.605	1.00	42.00	N
ANISOU	2722	NE2	GLN	B	106	5757	6121	4422	819	565	-415 N
ATOM	2723	N	ASP	B	107	57.555	-5.015	-2.974	1.00	41.36	N
ANISOU	2723	N	ASP	B	107	5360	5572	4775	709	727	-536 N
ATOM	2724	CA	ASP	B	107	58.907	-6.507	-2.719	1.00	42.45	C
ANISOU	2724	CA	ASP	B	107	5462	5674	4992	713	772	-582 C
ATOM	2725	C	ASP	B	107	53.921	-7.507	-1.501	1.00	41.30	C
ANISOU	2725	C	ASP	B	107	5299	5457	4070	688	6195	-633 C
ATOM	2726	O	ASP	B	107	59.594	-8.534	-1.701	1.00	42.73	O
ANISOU	2726	O	ASP	B	107	5458	5613	5155	710	691	-7131 O

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ATOM	2727	CB	ASP	B	107	59.862	-5.346	-2.443	1.00	45.98	C
ANISOU	2727	CB	ASP	B	107	5678	6114	5477	695	654	-505 C
ATOM	2728	CG	ASP	B	107	60.079	-4.520	-3.651	1.00	47.81	C
ANISOU	2728	CG	ASP	B	107	6138	6427	5601	723	353	-462 C
ATOM	2729	OD1	ASP	B	107	50.700	-5.022	-4.741	1.00	52.51	O
ANISOU	2729	OD1	ASP	B	107	6806	7127	6131	765	946	-509 O
ATOM	2730	OD2	ASP	B	107	60.617	-3.397	-8.560	1.00	45.25	O
ANISOU	2730	OD2	ASP	B	107	5501	6101	5292	704	1028	-383 O
ATOM	2731	N	THR	B	108	58.150	-7.244	-0.530	1.00	39.00	N
ANISOU	2731	N	THR	B	108	4993	5105	4720	646	633	-586 N
ATOM	2732	CA	THR	B	108	58.110	-3.163	0.593	1.30	42.70	C
ANISOU	2732	CA	THR	B	108	5436	5495	5291	619	559	-622 C
ATOM	2733	C	THR	B	108	57.502	-9.531	0.211	1.00	45.00	C
ANISOU	2733	C	THR	B	108	5755	5778	5565	634	435	-712 C
ATOM	2734	O	THR	B	108	58.052	-10.616	0.605	1.00	41.55	O
ANISOU	2734	O	THR	B	108	5312	5295	5216	640	457	-775 O
ATOM	2735	CB	THR	B	108	57.353	-7.542	1.816	1.00	36.64	C
ANISOU	2735	CB	THR	B	108	4663	4685	4573	570	512	-550 C
ATOM	2736	OG1	THR	B	108	58.087	-5.397	2.274	1.00	27.71	O
ANISOU	2736	OG1	THR	B	108	4772	4813	4743	553	577	-481 O
ATOM	2737	CG2	THR	B	108	57.244	-3.548	2.987	1.00	40.27	C
ANISOU	2737	CG2	THR	B	108	5103	5070	5126	542	434	-588 C
ATOM	2738	N	ALA	B	109	56.378	-3.496	-0.525	01.00	40.97	N
ANISOU	2738	N	ALA	B	109	5665	5701	5333	639	463	-718 N
ATOM	2739	CA	ALA	B	109	55.759	-10.739	-1.024	1.00	46.52	C
ANISOU	2739	CA	ALA	B	109	6010	6024	5635	649	385	-810 C
ATOM	2740	C	ALA	B	109	56.750	-11.583	-1.658	1.00	44.30	C
ANISOU	2740	C	ALA	B	109	5737	5751	5345	697	424	-903 C
ATOM	2741	O	ALA	B	109	56.750	-12.300	-1.710	1.00	49.52	O
ANISOU	2741	O	ALA	B	109	6402	6359	7055	693	372	-983 O
ATOM	2742	CB	ALA	B	109	54.504	-10.470	-1.337	1.00	42.46	C
ANISOU	2742	CB	ALA	B	109	5539	5584	5008	552	351	-804 C
ATOM	2743	N	LYS	B	110	57.605	-13.957	-2.682	1.00	45.38	N
ANISOU	2743	N3	LYS	B	110	5379	5948	5424	735	517	-391 N
ATOM	2744	CA	LYS	B	110	53.679	-11.699	-3.366	1.00	51.23	C
ANISOU	2744	CA	LYS	B	110	6508	6704	6171	733	575	-978 C
ATOM	2745	C	LYS	B	110	59.722	-12.293	-2.430	1.00	52.59	C
ANISOU	2745	C	LYS	B	110	6726	5785	5479	782	575	-999 C
ATOM	2746	O	LYS	B	110	60.204	-13.371	-2.767	1.00	55.93	O
ANISOU	2746	O	LYS	B	110	7143	7179	6922	815	557	-1089 O

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ATOM	2747	CB	LYS	B	110	59.419	-10.797	-4.335	1.00	59.22	C	
ANISOU	2747	CB	LYS	B	110	7612	7795	7092	813	675	-944	C
ATOM	2748	CG	LYS	B	110	58.679	-10.467	-5.614	1.00	59.42	C	
ANISOU	2748	CG	LYS	B	110	7639	7922	7957	844	636	-948	C
ATOM	2749	CD	LYS	B	110	59.555	-9.956	-6.671	1.00	67.57	C	
ANISOU	2749	CD	LYS	B	110	8719	9034	7920	889	301	-944	C
ATOM	2750	CE	LYS	B	110	50.224	-8.559	-6.359	1.30	72.77	C	
ANISOU	2750	CE	LYS	B	110	9351	9704	6596	888	833	-825	C
ATOM	2751	NZ	LYS	B	110	59.215	-7.455	-6.525	1.00	72.85	N	
ANISOU	2751	NZ	LYS	B	110	9399	9755	8530	850	874	-730	N
ATOM	2752	N	LEU	B	111	60.100	-11.564	-1.397	1.00	51.39	N	
ANISOU	2752	N	LEU	B	111	5498	5559	6355	749	583	-919	N
ATOM	2753	CA	LEU	B	111	51.177	-11.972	-0	437	1.30	51.30	C
ANISOU	2753	CA	LEU	B	111	5432	5479	5465	753	539	-929	C
ATOM	2754	C	LEU	B	111	63.402	-13.119	0.432	1.00	51.14	C	
ANISOU	2754	C	LEU	B	111	6453	4405	5573	733	492	-953	C
ATOM	2755	O	LEU	B	111	71.776	-13.880	0.853	1.00	43.97	O	
ANISOU	2755	O	LEU	B	111	5514	5443	5751	756	457	-1010	O
ATOM	2756	CB	LEU	B	111	51.577	-13.735	0.427	1.33	56.73	C	
ANISOU	2756	CB	LEU	B	111	7117	7194	7242	711	723	-433	C
ATOM	2757	CG	LEU	B	111	62.490	-9.522	0.045	1.30	53.15	C	
ANISOU	2757	CG	LEU	B	111	7265	7429	7399	714	729	-775	C
ATOM	2758	CD1	LEU	B	111	72.376	8-.559	1.345	1.00	50.45	C	
ANISOU	2758	CD1	LEU	B	111	6249	6411	5521	606	727	-733	C
ATOM	2759	CD2	LEU	B	111	63.730	-13.020	-0.795	1.00	55.15	C	
ANISOU	2759	CD2	LEU	B	111	6356	7089	7313	767	907	-938	C
ATOM	2760	N	ILE	B	112	59.527	-13.252	3.745	1.00	46.30	N	
ANISOU	2760	N	ILE	B	112	5939	5438	5006	735	417	-952	N
ATOM	2761	CA	ILE	B	112	59.083	-14.215	1.754	1.00	53.02	C	
ANISOU	2761	CA	ILE	B	112	6728	6535	6331	666	327	-969	C
ATOM	2762	C	ILE	B	112	59.569	-15.635	1.464	1.00	52.21	C	
ANISOU	2762	C	ILE	B	112	7734	5379	5823	705	302	-1071	C
ATOM	2763	O	ILE	B	112	59355	-15.157	0.377	1.00	47.53	O	
ANISOU	2763	O	ILE	B	112	5077	5318	6165	736	336	-1148	O
ATOM	2764	CB	ILE	B	112	57.526	-14.177	1.943	1.00	55.83	C	
ANISOU	2764	CB	ILE	B	112	7121	6894	7197	617	254	-943	C
ATOM	2765	CG1	ILE	B	112	57.123	-12.913	2.721	1.00	59.77	C	
ANISOU	2765	CG1	ILE	B	112	7602	7412	7695	575	263	-837	C
ATOM	2766	CG2	ILE	B	112	57.005	-14.426	2.653	1.00	50.60	C	
ANISOU	2766	CG2	ILE	B	112	7471	6145	6611	593	163	-382	C

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ATOM	2767	CD1	ILE	B	112	55.526	-12.617	2.709	1.00	59.89	C
ANISOU	2767	CD1	ILE	B	112	7646	7457	7653	539	211	-805 C
ATOM	2768	N	ASP	B	113	70.216	-16.260	2.455	1.00	55.06	N
ANISOU	2768	N	ASP	B	113	5955	6653	7298	737	275	-1072 N
ATOM	2769	CA	ASP	B	113	60.706	-17.638	2.292	1.00	54.52	C
ANISOU	2769	CA	ASP	B	113	6906	6522	7235	749	243	-1164 C
ATOM	2770	C	ASP	B	113	59.320	-18.543	2.709	1.00	54.94	C
ANISOU	2770	C	ASP	B	113	7000	6501	7375	707	151	-1180 C
ATOM	2771	O	ASP	B	113	59.327	-18.532	3.992	1.00	51.82	O
ANISOU	2771	O	ASP	B	113	5592	6051	7045	552	103	-1116 O
ATOM	2772	CB	ASP	B	113	61.962	-17.375	3.122	1.00	58.21	C
ANISOU	2772	CB	ASP	B	113	7295	6912	7836	777	263	-1152 C
ATOM	2773	CG	ASP	B	113	62.670	-19.161	2.737	1.00	66.49	C
ANISOU	2773	CG	ASP	B	113	3374	7929	8952	842	257	-1253 C
ATOM	2774	OD1	ASP	B	113	58.008	-19.140	2.453	1.00	68.45	O
ANISOU	2774	OD1	ASP	B	113	8577	8200	9233	899	321	-1279 O
ATOM	2775	OD2	ASP	B	113	61.997	-20.241	2.552	1.00	69.68	O
ANISOU	2775	OD2	ASP	B	113	8826	9257	9383	837	191	-1310 O
ATOM	2776	N	ARG	B	114	59.027	-19.302	1357	1.00	51.69	N
ANISOU	2776	N	ARG	B	114	6635	6080	5914	715	125	-1265 N
ATOM	2777	CA	ARG	B	114	57.861	-20.110	2.168	1.00	51.11	C
ANISOU	2777	CA	ARG	B	114	6604	5957	5860	669	87	-1287 C
ATOM	2778	C	ARG	B	114	55.174	-21.242	3.105	1.00	54.34	C
ANISOU	2778	C	ARG	B	114	7076	6305	7456	663	-18	-1301 C
ATOM	2779	O	ARG	B	114	57.279	-21.686	3.810	1.00	45.97	O
ANISOU	2779	O	ARG	B	114	6352	5503	5752	610	-89	-1278 O
ATOM	2780	CB	ARG	B	114	57.254	-20.721	0.885	1.00	51.45	C
ANISOU	2780	CB	ARG	B	114	6596	6030	6323	684	21	-1391 C
ATOM	2781	CG	ARG	B	114	58.132	-19.9041	0.245	3.00	55.56	C
ANISOU	2781	CG	ARG	B	114	7247	5563	7239	650	16	-1365 C
ATOM	2782	CD	ARG	B	114	56.889	-18.623	-0.378	1.00	62.75	C
ANISOU	2782	CD	ARG	B	114	8123	7670	8051	685	105	-1320 C
ATOM	2783	NE	ARG	B	114	55.603	-17.357	-1.150	1.00	63.47	N
ANISOU	2783	NE	ARG	B	114	8235	7862	8019	658	105	-1299 N
ATOM	2784	CZ	ARG	B	114	55.876	-16.596	-1.561	1.00	63.11	C
ANISOU	2784	CZ	ARG	B	114	8177	7908	7895	884	174	-1233 C
ATOM	2785	NH1	ARG	B	114	57.012	-15.043	-1.240	1.00	61.25	N
ANISOU	2785	NH1	ARG	B	114	7903	7674	7594	708	245	-1184 N
ATOM	2786	NH2	ARG	B	114	54.911	-15.955	-2.250	1.00	56.75	N
ANISOU	2786	NH2	ARG	B	114	7393	7189	6979	578	166	-1212 N

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ATOM	2787	N	LYS	B	115	59.400	-21.766	3.032	1.00	53	20	N
ANISOU	2787	N	LYS	B	115	6849	6055	7309	731	13	-1344	N
ATOM	2788	CA	LYS	B	115	59.583	-22.912	3.950	1.00	52	92	C
ANISOU	2788	CA	LYS	B	115	6819	5895	7394	737	-44	-1356	C
ATOM	2789	C	LYS	B	115	59.865	-22.461	5.410	1.00	45	53	C
ANISOU	2789	C	LYS	B	115	5973	5053	6655	702	-85	-1245	C
ATOM	2790	O	LYS	B	115	50.312	-23.076	6.305	1.00	45	47	O
ANISOU	2790	O	LYS	B	115	5855	4839	6583	659	-132	-1212	O
ATOM	2791	CB	LYS	B	115	60.892	-23.686	3.438	1.00	55	44	C
ANISOU	2791	CB	LYS	B	115	7129	8174	7750	823	-10	-1441	C
ATOM	2792	CG	LYS	B	115	60.648	-24.349	2.150	1.00	56	10	C
ANISOU	2792	CG	LYS	B	115	7262	8271	7754	357	-2	-1563	C
ATOM	2793	CD	LYS	B	115	61.785	-25.323	1.306	1.00	63	32	C
ANISOU	2793	CD	LYS	B	115	3173	7124	8763	946	21	-1555	C
ATOM	2794	CE	LYS	B	115	61.933	-26.431	2.628	1.00	64	27	C
ANISOU	2794	CE	LYS	B	115	3303	7098	9018	951	-46	-1849	C
ATOM	2795	NZ	LYS	B	115	61.058	-27.567	2.540	1.00	58	23	N
ANISOU	2795	NZ	LYS	B	115	8877	7510	9533	923	-115	-1725	N
ATOM	2796	N	SER	B	116	60.639	-21.435	5.536	1.00	43	95	N
ANISOU	2796	N	SER	B	116	6224	5425	5948	718	-8	-1139	N
ATOM	2797	CA	SER	B	116	60.700	-20.864	6.959	1.00	43	48	C
ANISOU	2797	CA	SER	B	116	5406	4721	6304	679	-25	-1057	C
ATOM	2798	C	SER	B	116	59.3113	-20.323	7.396	1.80	45	07	C
ANISOU	2798	C	SER	B	116	5721	4944	6450	597	-551	-1022	C
ATOM	2799	O	SER	B	116	58.925	-20.508	8.557	1.00	42	83	O
ANISOU	2799	O	SER	B	116	5436	4609	6230	553	-118	-960	O
ATOM	2800	CB	SER	B	116	51.824	-19.365	7.042	1.00	44	55	C
ANISOU	2800	CB	SER	B	116	5569	4921	6436	709	46	-1051	C
ATOM	2801	OG	SER	B	116	61.391	-18.654	6.571	1.00	50	14	O
ANISOU	2801	OG	SER	B	116	6275	5722	7055	680	92	-1015	O
ATOM	2802	N	LEU	B	117	58.536	-13.707	6.487	1.00	40	84	N
ANISOU	2802	N	LEU	B	117	5207	4486	5825	579	-43	-1036	N
ATOM	2803	CA	LEU	B	117	57.151	-19.365	6.866	1.00	41	66	C
ANISOU	2803	CA	LEU	B	117	5332	4605	5893	507	-39	-986	C
ATOM	2804	C	LEU	B	117	56.348	-30.516	7.343	1.00	43	85	C
ANISOU	2804	C	LEU	B	117	53.46	4791	6228	468	-172	-1007	C
ATOM	2805	O	LEU	B	117	55.680	-20.556	8.370	1.00	43	30	O
ANISOU	2805	O	LEU	B	117	5572	4693	6187	411	-216	-939	O
ATOM	2806	CB	LEU	B	117	56.375	-13.712	5.727	1.00	39	58	C
ANISOU	2806	CB	LEU	B	117	5083	4435	5515	501	-53	-1306	C

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ATOM	2807	CG	LEU	B	117	54.863	-18.532	5.971	1.00	40.55	C
ANISOU	2807	CG	LEU	B	117	5231	4578	5600	434	-115	-970 C
ATOM	2808	CD1	LEU	B	117	54.559	-17.534	7.149	1.00	41.59	C
ANISOU	2808	CD1	LEU	B	117	5332	4723	5748	391	-118	-384 C
ATOM	2809	CD2	LEU	B	117	54.216	-17.939	4.722	1.00	41.99	C
ANISOU	2809	CD2	LEU	B	117	5432	4858	5666	443	-92	-998 C
ATOM	2810	N	ALA	B	118	55349	-21.733	6.563	1.30	45.02	N
ANISOU	2810	N	ALA	B	118	5830	4892	6385	493	-152	-1102 N
ATOM	2811	CA	ALA	B	118	55.659	-22.505	7.010	1.00	44.30	C
ANISOU	2811	CA	ALA	B	118	5775	4697	5360	453	-260	-1125 C
ATOM	2812	C	ALA	B	118	56.140	-23.465	8.348	1.00	42.36	C
ANISOU	2812	C	ALA	B	118	5518	4355	6221	445	-302	-1066 C
ATOM	2813	O	ALA	B	118	55.304	-23.939	9.073	1.00	43.20	O
ANISOU	2813	O	ALA	B	118	5642	4409	5254	384	-350	-1029 O
ATOM	2814	CB	ALA	B	118	55.723	-24.073	5.954	1.00	46.50	C
ANISOU	2814	CB	ALA	B	118	6110	4941	6655	486	-283	-1248 C
ATOM	2815	N	ARG	B	119	57.450	-23.411	8.680	1.00	41.52	N
ANISOU	2815	N	ARG	B	119	5391	4245	5178	507	-269	-1055 N
ATOM	2616	CA	ARG	B	119	57.900	-23.554	10.014	1.00	40.33	C
ANISOU	2816	CA	ARG	B	119	5181	3969	5076	503	-306	-990 C
ATOM	2817	C	ARG	B	119	57.357	-22.945	11.055	1.30	39.92	C
ANISOU	2817	C	ARG	B	119	5022	3878	5925	442	-346	-835 C
ATOM	2818	O	ARG	B	119	57.030	-23.065	12.210	1.03	38.52	O
ANISOU	2818	O	ARG	B	119	4966	3752	5915	431	-368	-821 O
ATOM	2819	CB	ARG	B	119	59.425	-23.831	10.158	1.00	43.47	C
ANISOU	2819	CB	ARG	B	119	5568	4391	6557	582	-268	-937 C
ATOM	2820	CG	ARG	B	119	60.140	-24.949	9.451	1.00	47.87	C
ANISOU	2820	CG	ARG	B	119	6146	4876	7150	354	-256	-1094 C
ATOM	2821	CD	ARG	B	119	51.562	-25.129	10.012	1.00	49.15	C
ANISOU	2821	CD	ARG	B	119	6262	5011	7404	726	-250	-1079 C
ATOM	2822	NE	ARG	B	119	60.370	-23.921	9.901	1.00	46.67	N
ANISOU	2822	NE	ARG	B	119	5909	4828	7073	750	-182	-1053 N
ATOM	2823	CZ	ARG	B	119	63.024	-23.573	8.803	1.00	55.78	C
ANISOU	2823	CZ	ARG	B	119	7015	5023	8154	802	-112	-1717 C
ATOM	2824	NH1	ARG	B	119	62.552	-24.318	7.710	1.00	54.58	N
ANISOU	3824	NH1	ARG	B	119	5903	5849	7987	341	-102	-1219 N
ATOM	2825	NH2	ARG	B	119	63.750	-22.419	8.738	1.00	55.16	N
ANISOU	2825	NH2	ARG	B	119	6870	6037	8045	813	-49	-1333 N
ATOM	2826	N	ILE	B	120	57.287	-21.650	10.700	1.00	37.34	N
ANISOU	2826	N	ILE	B	120	4751	3773	5523	426	-264	-861 N

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ATOM	2827	CA	ILE	B	120	55.856	-20.556	11.599	1.00	36.56	C
ANISOU	2827	CA	ILE	B	120	4632	3596	5391	355	-266	-754 C
ATOM	2828	C	ILE	B	120	55.363	-20.889	12.035	1.30	36.22	C
ANISOU	2828	C	ILE	B	120	4644	3667	5452	309	-318	-737 C
ATOM	2829	O	ILE	B	120	64.929	-20.535	13.146	1.00	36.19	O
ANISOU	2829	O	ILE	B	120	4533	3043	5474	263	-351	-665 O
ATOM	2830	CB	ILE	B	120	57.133	-19.209	11.291	1.00	35.05	C
ANISOU	2830	CB	ILE	B	120	4432	3684	5253	397	-195	-743 C
ATOM	2831	CG1	ILE	B	120	58.655	-18.918	11.294	1.00	35.24	C
ANISOU	2831	CG1	ILE	B	120	4536	3794	5437	457	-149	-751 C
ATOM	2832	CG2	ILE	B	120	56.371	-18.223	32.255	1.00	37.19	C
ANISOU	2832	CG2	ILE	B	120	4557	3946	5496	1336	-207	-652 C
ATOM	2833	CD1	ILE	B	120	59.016	-17.506	10.545	1.00	34.67	C
ANISOU	2833	CD1	ILE	B	120	4313	3694	5367	474	-70	-748 C
ATOM	2834	N	VAL	B	121	54.501	-21.204	10.970	1.00	35.55	N
ANISOU	2834	N	VAL	B	121	4503	3610	5884	208	-325	-581 N
ATOM	2835	CA	VAL	B	121	53.165	-21.532	11.136	1.00	35.54	C
ANISOU	2835	CA	VAL	B	121	4512	3585	5333	225	-377	-758 C
ATOM	2886	C	VAL	B	121	58.019	-22.693	12.125	1.00	36.21	C
ANISOU	2886	C	VAL	B	121	4717	3561	5481	191	-437	-763 C
ATOM	2837	O	VAL	B	121	52.233	-22.001	13.057	1.00	37.19	O
ANISOU	2837	O	VAL	B	121	4635	3581	5513	129	-467	-693 O
ATOM	2838	CB	VAL	B	121	52.523	-21.787	9.765	3.00	35.68	C
ANISOU	2838	CB	VAL	B	121	4657	3639	5259	220	-379	-874 C
ATOM	2839	CG1	VAL	B	121	51.295	-22.675	9.842	1.00	35.99	C
ANISOU	2839	CG1	VAL	B	121	4852	3759	5448	157	-444	-893 C
ATOM	2840	CG2	VAL	B	121	52.233	-20.486	9.085	1.00	34.27	C
ANISOU	2840	CG2	VAL	B	121	4460	3581	4080	237	-831	-861 C
ATOM	2841	N	GLU	B	122	53.620	-23.755	11.088	1.00	37.04	N
ANISOU	2841	N	GLU	B	122	4844	3574	5556	234	-451	-833 N
ATOM	2842	CA	GLU	B	122	53.731	-24.910	12.889	1.00	39.33	C
ANISOU	2842	CA	GLU	B	122	5100	3745	6037	207	-537	-755 C
ATOM	2843	C	GLU	B	122	24.173	-24.560	14.277	1.00	40.12	C
ANISOU	2843	C	GLU	B	122	5231	3838	6174	202	-513	-685 C
ATOM	2844	O	GLU	B	122	53.607	-25.051	15.243	1.00	39.08	O
ANISOU	2844	O	GLU	B	122	5113	3553	6082	148	-557	-623 O
ATOM	2845	CB	GLU	B	122	54.579	-25.157	12.417	1.00	41.21	C
ANISOU	2845	CB	GLU	B	122	5431	3877	6359	268	-520	-866 C
ATOM	2846	CG	GLU	B	122	54.317	-25.755	11.135	1.00	53.56	C
ANISOU	2846	CG	GLU	B	122	7035	5428	7588	263	-529	-975 C

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ATOM	2847	CD	GLU	B	122	52.530	-27.064	13.037	1.00	56.80	C	
ANISOU	2847	CD	GLU	B	122	7457	5827	5257	155	-578	-958 C	
ATOM	2848	OE1	GLU	B	122	52.216	-27.743	12.337	1.00	59.54	O	
ANISOU	2848	OE1	GLU	B	122	9097	7354	9972	117	-523	-898 O	
ATOM	2849	OE2	GLU	B	122	51.722	-26.588	10.457	1.00	59.44	O	
ANISOU	2849	OE2	GLU	B	122	7500	6245	8540	137	-571	-909 O	
ATOM	2850	N	ARG	B	123	55.196	-23.724	14.399	1.00	40.06	N	
ANISOU	2850	N	ARG	B	123	5184	3655	6152	255	-468	-667 N	
ATOM	2851	CA	ARG	B	123	55.555	-23.277	15.736	1.00	89.93	C	
ANISOU	2851	CA	ARG	B	123	5137	3376	5159	245	-476	-573 C	
ATOM	2852	C	ARG	B	123	54.460	-22.535	10.452	1.00	30.82	C	
ANISOU	2852	C	ARG	B	123	5113	3920	6099	170	-485	-499 C	
ATOM	2853	O	ARG	B	123	64.212	-22.743	17.644	1.00	85.79	O	
ANISOU	2853	O	ARG	B	123	4730	3506	5742	132	-519	-425 O	
ATOM	2854	CB	ARG	B	123	55.777	-22.380	15.027	3.33	45.15	C	
ANISOU	2854	CB	ARG	B	123	5752	4500	6808	306	-423	-575 C	
ATOM	2855	CG	ARG	B	123	57.963	-23.154	15.051	1.30	45.54	C	
ANISOU	2855	CG	ARG	B	123	5926	4715	7339	307	-414	-643 C	
ATOM	2856	CD	ARG	B	123	59.233	-22.733	15.711	1.00	51.95	C	
ANISOU	2856	CD	ARG	B	123	6559	5424	7717	436	-395	-608 C	
ATOM	2857	NE	ARG	B	123	59.939	-23.858	13.263	1.00	52.03	N	
ANISOU	2857	NE	ARG	B	123	7843	6600	9125	482	-438	-605 N	
ATOM	2858	CZ	ARG	B	123	61.054	-24.410	15.728	1.00	61.73	C	
ANISOU	2858	CZ	ARG	B	123	7798	6534	9141	5513	-422	-666 C	
ATOM	2859	NH	ARG	B	123	1	61.556	-23.967	14.576	1.00	52.06	N
ANISOU	2859	NH1	ARG	B	123	6546	5331	7872	605	-351	-739 N	
ATOM	2860	NH2	ARG	B	123	81.659	-25.428	16.350	1.00	70.92	N	
ANISOU	2860	NH2	ARG	B	123	8953	7599	10385	605	-456	-552 N	
ATOM	2861	N	VAL	B	124	53.840	-21.592	15.759	1.03	33.45	N	
ANISOU	2861	N	VAL	B	124	4297	3234	5212	153	-450	-516 N	
ATOM	2862	CA	VAL	B	124	52.704	-20.603	16.396	1.00	37.09	C	
ANISOU	2862	CA	VAL	B	124	4746	3729	5325	85	-458	-453 C	
ATOM	2863	C	VAL	B	124	51.559	-21.870	16.724	1.00	34.82	C	
ANISOU	2863	C	VAL	B	124	4489	3377	5363	18	-512	-438 C	
ATOM	2864	O	VAL	B	124	50.969	-21.826	17.812	1.00	36.38	O	
ANISOU	2864	O	VAL	B	124	4680	3575	5567	-35	-535	-363 O	
ATOM	2865	CB	VAL	B	124	52.162	-19.789	15.475	1.09	37.47	C	
ANISOU	2865	CB	VAL	B	124	4780	3870	5585	85	-415	-475 C	
ATOM	2866	CG1	VAL	B	124	50.940	-19.154	16.125	1.00	39.32	C	
ANISOU	2866	CG1	VAL	B	124	5003	4160	5778	21	-425	-413 C	

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ATOM	2867	CG2	VAL	B	124	53.266	-18.779	15.165	1.00	34.02	C
ANISOU	2867	CG2	VAL	B	124	4315	3487	5125	144	-353	-481 C
ATOM	2868	N	HIS	B	125	51.262	-22.763	15.782	1.30	37.17	N
ANISOU	2868	N	HIS	B	125	4907	3715	3764	15	-532	-512 N
ATOM	2869	CA	HIS	B	125	50.179	-23.738	16.014	1.00	38.87	C
ANISOU	2869	CA	HIS	B	125	5937	3758	5807	-55	-584	-507 C
ATOM	2870	C	HIS	B	125	50.464	-24.517	17.292	1.00	40.10	C
ANISOU	2870	C	HIS	B	125	5231	3843	6356	-75	-619	-437 C
ATOM	2871	O	HIS	B	125	49.543	-24.721	18.117	1.00	36.50	O
ANISOU	2871	O	HIS	B	125	4778	3284	5707	-148	-645	-373 O
ATOM	2872	CB	HIS	B	125	50.008	-24.668	14.840	1.00	38.04	C
ANISOU	2872	CB	HIS	B	125	4993	3426	5830	-50	-602	-607 C
ATOM	2873	CG	HIS	B	125	49.212	-25.340	15.134	1.00	43.94	C
ANISOU	2873	CG	HIS	B	125	5774	4285	6536	-123	-858	-605 C
ATOM	2874	ND1	HIS	B	125	47.799	-25.605	15.385	1.00	43.16	N
ANISOU	2874	ND1	HIS	B	125	5063	4228	6505	-206	-678	-573 N
ATOM	2875	CD2	HIS	B	125	49.380	-27.205	15.214	1.00	44.24	C
ANISOU	2875	CD2	HIS	B	125	5857	4191	6763	-128	-466	-534 C
ATOM	2876	CE1	HIS	B	125	47.254	-26.915	15.005	1.00	45.58	C
ANISOU	2876	CE1	HIS	B	125	5.002	4433	5482	-256	-726	-579 C
ATOM	2877	NE2	HIS	B	125	48.215	-27.824	15.493	1.00	43.63	N
ANISOU	2877	NE2	HIS	B	125	5794	4077	0707	-210	-737	-517 N
ATOM	2878	N	GLN	B	126	51.734	-24.214	17.495	1.00	38.20	N
ANISOU	2878	N	GLN	B	126	4997	3545	5970	-9	-517	-442 N
ATOM	2879	CA	GLN	B	126	52.050	-25.737	18.701	1.00	43.11	C
ANISOU	2879	CA	GLN	B	126	5837	4077	8667	-13	-656	-372 C
ATOM	2880	C	GLN	B	126	52.166	-24.918	19.950	1.00	41.20	C
ANISOU	2880	C	GLN	B	126	5051	3893	6400	-32	-549	-274 C
ATOM	2881	O	GLN	B	126	51.873	-25.440	21.015	1.00	40.48	O
ANISOU	2881	O	GLN	B	126	5284	3755	5042	-73	-683	-200 O
ATOM	2882	CB	GLN	B	126	53.384	-26.548	8.558	1.20	47.91	C
ANISOU	2882	CB	GLN	B	126	6261	4592	7350	64	-665	-407 C
ATOM	2883	CG	GLN	B	126	53.542	-27.363	17.296	1.00	57.95	C
ANISOU	2883	CG	GLN	B	126	7567	5892	6550	98	-566	-518 C
ATOM	2884	CD	GLN	B	126	62.454	-24.387	17.061	1.00	72.90	C
ANISOU	2884	CD	GLN	B	126	9509	7616	19575	25	-708	-542 C
ATOM	2885	OE1	GLN	B	126	52.121	-29.185	17.950	1.00	82.84	O
ANISOU	2885	OE1	GLN	B	126	10693	6586	11792	-16	-750	-430 O
ATOM	2885	NE2	GLN	B	126	51.904	-28.397	15.833	1.00	75.38	N
ANISOU	2886	NE2	GLN	B	126	9834	7958	10953	16	-699	-635 N

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ATOM	2887	N	ALA	B	127	52.559	-23.629	19.851	1.00	85.97	N
ANISOU	2887	N	ALA	B	127	4657	3334	5677	-3	-604	-278 N
ATOM	2888	CA	ALA	B	127	52.754	-22.775	21.055	1.30	38.96	C
ANISOU	2888	CA	ALA	B	127	5035	3772	6032	-12	-595	-339 C
ATOM	2889	C	ALA	B	127	51.522	-22.702	21.989	1.30	37.39	C
ANISOU	2889	C	ALA	B	127	4807	3595	5805	-95	-516	-114 C
ATOM	2890	O	ALA	B	127	50.336	-22.522	21.575	1.00	37.95	O
ANISOU	2890	O	ALA	B	127	4880	3704	5837	-150	-612	-326 O
ATOM	2891	CB	ALA	B	127	53.256	-21.344	23.657	3.03	32.20	C
ANISOU	2891	CB	ALA	B	127	4231	3144	5240	2	-54	-21 C
ATOM	2892	N	GLU	B	128	51.799	-22.334	23.271	1.30	38.60	N
ANISOU	2892	N	GLU	B	128	4958	3737	5977	-105	-538	-35 N
ATOM	2893	CA	GLU	B	128	50.738	-22.505	24.251	1.00	37.66	C
ANISOU	2893	CA	GLU	B	128	4883	3559	5318	-174	-544	40 C
ATOM	2894	C	GLU	B	128	50.743	-21.930	24.664	1.90	87.30	C
ANISOU	2894	C	GLU	B	128	4744	3727	5700	-173	-605	62 C
ATOM	2895	O	GLU	B	128	4.5157	-29.525	25.353	1.00	34.75	O
ANISOU	2895	O	GLU	B	128	4409	3455	5325	-226	-595	96 O
ATOM	2896	CB	GLU	B	128	51.036	-23.382	25.495	1.00	39.70	C
ANISOU	2896	CB	GLU	B	128	5113	3860	6123	-199	-687	121 C
ATOM	2897	CG	GLU	B	128	50.151	-24.657	25.314	1.00	50.78	C
ANISOU	2897	CG	GLU	B	128	5558	5161	7575	-247	-720	126 C
ATOM	2898	CD	GLU	B	128	52.772	-25.652	25.961	1.00	51.42	C
ANISOU	2898	CD	GLU	B	128	6677	5128	7731	-233	-763	172 C
ATOM	2899	OE1	GLU	B	128	51.412	-25.707	27.039	1.00	62.05	O
ANISOU	2899	OE1	GLU	B	128	5316	6487	9375	-206	-776	241 O
ATOM	2900	OE2	GLU	B	128	50.584	-27.031	25.463	1.00	64.06	O
ANISOU	2900	OE2	GLU	B	128	8320	6623	9397	-243	-790	144 O
ATOM	2901	N	PHE	B	129	51.396	-20.343	24.537	1.00	31.52	N
ANISOU	2901	N	PHE	B	129	4001	3035	4977	-111	-533	40 N
ATOM	2902	CA	PHE	B	129	51.997	-16.973	25.058	1.00	31.55	C
ANISOU	2902	CA	PHE	B	129	3957	3121	4930	-103	-549	60 C
ATOM	2903	C	PHE	B	129	52.963	-18.254	24.090	1.00	36.67	C
ANISOU	2903	C	PHE	B	129	4533	3795	5554	-47	-509	-6 C
ATOM	2904	O	PHE	B	129	54.067	-15.787	23.797	1.00	33.20	O
ANISOU	2904	O	PHE	B	129	4142	3335	5165	5	-537	-35 O
ATOM	2905	CB	PHE	B	129	52.591	-18.942	26.455	1.00	34.58	C
ANISOU	2905	CB	PHE	B	129	4331	3509	5300	-105	-572	127 C
ATOM	2906	CG	PHE	B	129	52.374	-17.545	26.913	1.90	36.55	C
ANISOU	2906	CG	PHE	B	129	4545	3846	5497	-98	-539	134 C

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ATOM	2907	CD1	PHE	B	129	51.834	-36.799	27.515	1.30	37.98	C
ANISOU	2907	CD1	PHE	B	129	4605	8981	5503	-147	-523	170 C
ATOM	2908	CD2	PHE	B	129	54.138	-16.950	26.693	1.00	42.04	C
ANISOU	2908	CD2	PHE	B	129	5212	4557	6205	-40	-520	99 C
ATOM	2909	CE1	PHE	B	129	62.076	-15.495	27.956	3.00	44.37	C
ANISOU	2909	CE1	PHE	B	129	5563	5041	5443	-139	-492	171 C
ATOM	2910	CE2	PHE	B	129	54.351	-15.637	27.093	1.00	52.20	C
ANISOU	2910	CE2	PHE	B	129	5213	4563	6192	-44	-488	101 C
ATOM	2911	CZ	PHE	B	129	53.335	-14.924	27.723	1.03	39.44	C
ANISOU	2911	CZ	PHE	B	129	4852	4361	3771	-91	-475	336 C
ATOM	2912	N	ILE	B	130	52.566	-17.105	22.551	1.00	30.00	N
ANISOU	2912	N	ILE	B	130	3721	3025	4654	-50	-4154	-29 N
ATOM	2913	CA	ILE	B	130	53.418	-15.364	22.652	1.00	30.11	C
ANISOU	2913	CA	ILE	B	130	3715	3055	4659	12	-420	-82 C
ATOM	2914	C	ILE	B	130	53.721	-14.1988	23.279	1.00	28.07	C
ANISOU	2914	C	ILE	B	130	3425	2877	4362	0	-387	-57 C
ATOM	2915	O	ILE	B	130	52.805	-14.206	23.611	1.00	31.15	O
ANISOU	2915	O	ILE	B	130	3812	3320	4702	-85	-372	-31 O
ATOM	2916	CB	ILE	B	130	52.668	-16.168	21.279	1.30	28.93	C
ANISOU	2916	CB	ILE	B	130	3578	2939	4473	-1	-393	-135 C
ATOM	2917	CG1	ILE	B	130	52.387	-17.534	20.731	1.00	30.31	C
ANISOU	2917	CG1	ILE	B	130	3766	3041	4688	-8	-430	-165 C
ATOM	2918	CG2	ILE	B	130	53.529	-15.405	20.340	1.00	30.91	C
ANISOU	2918	CG2	ILE	B	130	38.13	3223	4719	53	-342	-192 C
ATOM	2919	CD1	ILE	B	130	51.497	-17.599	19.462	1.00	33.21	C
ANISOU	2919	CD1	ILE	B	130	4168	3431	5018	-16	-418	-219 C
ATOM	2920	N	GLY	B	131	54.965	-14.666	23.391	1.00	28.55	N
ANISOU	2920	N	GLY	B	131	3461	2939	4449	39	-374	-70 N
ATOM	2921	CA	GLY	B	131	55.419	-13.354	23.871	1.00	29.09	C
ANISOU	2921	CA	GLY	B	131	3496	3065	4489	89	-341	-59 C
ATOM	2922	C	GLY	B	131	55.960	-12.593	22.696	1.00	32.60	C
ANISOU	2922	C	GLY	B	131	3929	3534	4924	72	-284	-109 C
ATOM	2923	O	GLY	B	131	55.649	-13.134	21.600	1.00	29.74	O
ANISOU	2923	O	GLY	B	131	3565	3141	4594	111	-275	-153 O
ATOM	2924	N	CYS	B	132	55.630	-11.299	22.656	1.00	29.04	N
ANISOU	2924	N	CYS	B	132	3470	4137	4428	57	-241	-134 N
ATOM	2925	CA	CYS	B	132	53.192	-10.445	21.586	1.00	29.01	C
ANISOU	2925	CA	CYS	B	132	3453	3155	4413	86	-182	-141 C
ATOM	2926	C	CYS	B	132	50.875	-9.255	22.224	1.00	26.62	C
ANISOU	2926	C	CYS	B	132	3120	2895	4109	78	-152	-130 C

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ATOM	2927	O	CYS	B	132	56.465	-3.759	23.321	1.00	27.67	O	
ANISOU	2927	O	CYS	B	132	3277	3063	4249	46	-168	-95	O
ATOM	2928	CB	CYS	B	132	53.373	-10.038	23.806	1.00	29.98	C	
ANISOU	2928	CB	CYS	B	132	3693	3396	4481	90	-151	-150	C
ATOM	2929	SG	CYS	B	132	53.909	-11.259	23.064	1.00	32.19	S	
ANISOU	2929	SG	CYS	B	132	3916	3563	4751	67	-194	-150	S
ATOM	2930	N	ILE	B	133	57.915	-8.781	71.537	1.09	29.73	N	
ANISOU	2930	N	ILE	B	133	3489	3265	4521	105	-197	-161	N
ATOM	2931	CA	ILE	B	133	56.772	-7.732	22.096	1.00	29.79	C	
ANISOU	2931	CA	ILE	B	133	3462	3315	4542	95	-81	-159	C
ATOM	2932	C	ILE	B	133	58.041	-5.757	20.972	1.00	21.06	C	
ANISOU	2932	C	ILE	B	133	3520	3495	4585	107	-8	-160	C
ATOM	2933	O	ILE	B	133	59.379	-7.189	19.359	1.00	30.27	O	
ANISOU	2933	O	ILE	B	133	3521	3389	4590	139	17	-209	O
ATOM	2934	CB	ILE	B	133	50.105	-5.319	22.507	1.00	33.25	C	
ANISOU	2934	CB	ILE	B	133	3858	3737	5040	116	-106	-173	C
ATOM	2935	CG1	ILE	B	133	59.944	-9.575	23.368	1.00	34.32	C	
ANISOU	2935	CG2	ILE	B	133	4002	3841	5199	116	-163	-151	C
ATOM	2936	CG2	ILE	B	133	60.656	-7.281	23.338	1.00	37.59	C	
ANISOU	2936	CG2	ILE	B	133	4368	4314	5601	94	-94	-169	C
ATOM	2937	CD1	ILE	B	133	61.248	-19.440	23.456	1.00	96.33	C	
ANISOU	2937	CD1	ILE	B	133	4218	4073	5517	159	-207	-170	C
ATOM	2938	N	GLY	B	134	59.018	-5.434	21.219	1.00	20.10	N	
ANISOU	2938	N	GLY	B	134	3405	3397	4546	84	33	-163	N
ATOM	2939	CA	GLY	B	134	59.311	-4.496	20.174	1.06	29.82	C	
ANISOU	2939	CA	GLY	B	134	3459	3374	4496	84	104	-160	C
ATOM	2940	C	GLY	B	134	59.100	-3.100	20.746	1.00	30.79	C	
ANISOU	2940	C	GLY	B	134	3586	3509	4605	52	133	-151	C
ATOM	2941	O	GLY	B	134	158.291	-2.961	21.655	1.30	30.95	O	
ANISOU	2941	O	GLY	B	134	3621	3533	4604	41	100	-140	O
ATOM	2942	N	VAL	B	135	59.902	-2.122	20.230	1.90	29.31	N	
ANISOU	2942	N	VAL	B	135	4254	3197	4307	57	1134	-170	N
ATOM	2943	CA	VAL	B	135	59.819	-0.722	20.850	1.00	29.80	C	
ANISOU	2943	CA	VAL	B	135	3446	3384	4491	24	226	-157	C
ATOM	2944	C	VAL	B	135	59.664	0.307	19.805	1.00	32.21	C	
ANISOU	2944	C	VAL	B	135	3777	3686	4774	30	300	-145	O
ATOM	2945	O	VAL	B	135	59.803	-0.018	18.126	1.00	25.82	O	
ANISOU	2945	O	VAL	B	135	3481	3393	4457	59	332	-149	C
ATOM	2946	CB	VAL	B	135	51.037	-0.037	21.740	1.00	34.09	C	
ANISOU	2946	CB	VAL	B	135	3939	3925	5089	-7	220	-176	C

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ATOM	2947	CG1	VAL	B	135	61.116	-1.323	22.950	1.00	29.80	C	
ANISOU	2947	CG1	VAL	B	135	3374	3387	4560	-11	138	-179	C
ATOM	2948	CG2	VAL	B	135	62.338	-0.343	20.924	1.00	26.40	C	
ANISOU	2948	CG2	VAL	B	135	2921	2952	4157	2	255	-199	C
ATOM	2949	N	GLY	B	136	59.231	1.542	20.192	1.00	29.65	N	
ANISOU	2949	N	GLY	B	136	3472	3354	4439	7	328	-129	N
ATOM	2950	CA	GLY	B	136	59.134	2.552	19.002	1.00	30.39	C	
ANISOU	2950	CA	GLY	B	136	3557	3502	4576	18	404	-110	C
ATOM	2951	C	GLY	B	136	58.205	2.147	13.022	1.00	28.81	C	
ANISOU	2951	C	GLY	B	136	3395	3220	4217	59	410	-91	C
ATOM	2952	O	GLY	B	136	57.117	1.576	18.362	1.00	29.15	O	
ANISOU	2952	O	GLY	B	136	3369	3188	4138	70	353	-84	O
ATOM	2953	N	ALA	B	137	58.514	2.382	16.716	1.00	31.76	N	
ANISOU	2953	N	ALA	B	137	3617	3638	4511	82	466	-84	N
ATOM	2954	CA	ALA	B	137	57.548	1.950	15.747	1.00	27.34	C	
ANISOU	2954	CA	ALA	B	137	3295	3104	3980	122	404	-70	C
ATOM	2955	C	ALA	B	137	57.328	0.428	15.521	1.00	30.37	C	
ANISOU	2955	C	ALA	B	137	3569	3506	4363	140	404	-93	C
ATOM	2956	O	ALA	B	137	56.235	0.001	15.197	1.00	24.55	O	
ANISOU	2956	O	ALA	B	137	2974	2803	3589	162	377	-91	O
ATOM	2957	CB	ALA	B	137	57.002	2.559	14.358	1.00	32.93	C	
ANISOU	2957	CB	ALA	B	137	4023	3323	4674	144	541	-82	C
ATOM	2958	N	SER	B	138	58.308	-0.388	16.004	1.00	27.28	N	
ANISOU	2958	N	SER	B	138	3237	3106	4021	132	381	-130	N
ATOM	2959	CA	SER	B	138	58.026	-1.821	16.061	1.00	27.03	C	
ANISOU	2959	CA	SER	B	138	3210	3086	3997	148	319	-154	C
ATOM	2960	C	SER	B	138	56.951	-2.227	17.086	1.00	28.22	C	
ANISOU	2960	C	SER	B	138	3365	3225	4131	130	251	-140	C
ATOM	2961	O	SER	B	138	55.413	-3.322	15.955	1.00	28.28	O	
ANISOU	2961	O	SER	B	138	3384	3234	4108	140	204	-152	O
ATOM	2962	CB	SER	B	138	59.311	-2.644	15.353	1.00	25.20	C	
ANISOU	2962	CB	SER	B	138	2924	2033	3819	149	303	-138	C
ATOM	2963	OG	SER	B	138	60.159	-2.541	15.176	1.00	27.70	O	
ANISOU	2963	OG	SER	B	138	3228	3163	4133	174	366	-207	O
ATOM	2964	N	SER	B	139	56.579	-1.392	18.080	1.30	29.89	N	
ANISOU	2964	N	SER	B	139	3573	3431	4349	102	243	-119	N
ATOM	2965	CA	SER	B	139	55.573	-1.660	19.032	1.50	27.45	C	
ANISOU	2965	CA	SER	B	139	3281	3127	4022	86	194	-102	C
ATOM	2966	C	SER	B	139	54.220	-1.785	18.268	1.00	31.03	C	
ANISOU	2966	C	SER	B	139	3758	3605	4415	1051	189	-87	C

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ATOM	2967	O	SER	B	139	53.395	-2.610	18.592	1.00	29.38	O
ANISOU	2967	O	SER	B	139	3554	3405	4194	130	138	-85
ATOM	2968	CB	SER	B	139	55.468	-0.539	25.034	1.00	32.62	C
ANISOU	2968	CB	SER	B	139	3925	3778	4582	50	205	-85
ATOM	2969	OG	SER	B	139	54.285	-0.564	20.820	1.00	27.68	O
ANISOU	2969	OG	SER	B	139	4587	4431	5292	49	166	-67
ATOM	2970	N	ILE	B	140	54.056	-1.021	17.189	3.00	29.53	N
ANISOU	2970	N	ILE	B	140	3613	3442	4204	131	241	-77
ATOM	2971	CA	ILE	B	140	52.809	-1.070	15.318	1.00	32.52	C
ANISOU	2971	CA	ILE	B	140	4908	2841	4539	157	238	-64
ATOM	2972	C	ILE	B	140	52.567	-2.458	15.659	1.00	31.52	C
ANISOU	2972	C	ILE	B	140	3882	3724	4372	169	197	-95
ATOM	2973	O	ILE	B	140	51.569	-3.043	15.029	1.00	27.97	O
ANISOU	2973	O	ILE	B	140	3440	3294	3893	156	153	-94
ATOM	2974	CB	ILE	B	140	52.889	0.058	15.257	1.00	82.72	C
ANISOU	2974	CB	ILE	B	140	4057	3878	4497	186	303	-42
ATOM	2975	CG1	ILE	B	140	52.403	1.376	15.905	1.00	83.96	C
ANISOU	2975	CG1	ILE	B	140	3847	3345	4273	180	329	-9
ATOM	2976	CG2	ILE	B	140	52.373	-0.285	13.940	1.00	37.75	C
ANISOU	2976	CG2	ILE	B	140	4719	4555	5069	222	302	-42
ATOM	2977	CD1	ILE	B	140	521.377	1.935	16.861	1.00	33.86	C
ANISOU	2977	CD1	ILE	B	140	4195	3974	4607	147	345	-15
ATOM	2978	N	VAL	B	141	53.777	-2.950	15.097	1.00	28.10	N
ANISOU	2978	N	VAL	B	141	3438	3277	3953	181	215	-125
ATOM	2979	CA	VAL	B	141	53.826	-4.256	14	504	1.00	27.44
ANISOU	2979	CA	VAL	B	141	3355	3191	3879	194	162	-153
ATOM	2980	C	VAL	B	141	53.633	-5.392	15.499	1.00	26.34	C
ANISOU	2980	C	VAL	B	141	3204	3025	3779	168	112	-173
ATOM	2981	O	VAL	B	141	52.819	-6.304	15.208	1.00	29.80	O
ANISOU	2981	O	VAL	B	141	2653	2467	4201	163	69	-188
ATOM	2982	CB	VAL	B	141	55.128	-4.432	13.699	1.00	26.68	C
ANISOU	2982	CB	VAL	B	141	3247	3086	3801	218	225	-194
ATOM	2983	CG1	VAL	B	141	55.190	-5.838	13.122	1.00	28.12	C
ANISOU	2983	CG1	VAL	B	141	3434	3263	3987	236	189	-242
ATOM	2984	CG2	VAL	B	141	55.228	-3.236	12.723	1.00	28.41	C
ANISOU	2984	CG2	VAL	B	141	3484	3337	3975	240	298	-172
ATOM	2985	N	GLY	B	142	54.234	-5.326	13.663	1.00	26.80	N
ANISOU	2985	N	GLY	B	142	3238	3057	3388	145	100	-162
ATOM	2986	CA	GLY	B	142	54.110	-6.400	17.554	1.00	25.41	C
ANISOU	2986	CA	GLY	B	142	3052	2852	3748	121	34	-162

TABLE 3-continued

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ATOM	2987	C	GLY	B	142	52.656	-6.394	18.250	1.00	25.75	C
ANISOU	2987	C	GLY	B	142	3110	2016	3759	94	-2	-132 C
ATOM	2988	O	GLY	B	142	52.025	-7.443	18.495	1.00	24.56	O
ANISOU	2988	O	GLY	B	142	2065	2753	3613	77	-53	-134 O
ATOM	2989	N	ARG	B	143	52.034	-5.196	18.453	1.00	24.67	N
ANISOU	2989	N	ARG	B	143	2977	2808	3588	90	28	-104 N
ATOM	2990	CA	ARG	B	143	50.635	-5.090	18.817	1.00	26.47	C
ANISOU	2990	CA	ARG	B	143	3213	3065	3780	73	5	-70 C
ATOM	2991	C	ARG	B	143	49.745	-5.677	17.782	1.00	22.37	C
ANISOU	2991	C	ARG	B	143	2771	2333	3286	84	-13	-93 C
ATOM	2992	O	ARG	B	143	48.752	-6.387	18.106	1.00	24.30	O
ANISOU	2992	O	ARG	B	143	2905	2869	3505	59	-58	-86 O
ATOM	2993	CB	ARG	B	143	50.271	-3.637	19.155	1.00	26.92	C
ANISOU	2993	CB	ARG	B	143	3272	3146	3839	77	44	-51 C
ATOM	2994	CG	ARG	B	143	50.976	-3	059	20.395	1.00	32.67 C
ANISOU	2994	CG	ARG	B	143	3986	3857	4571	56	52	-41 C
ATOM	2995	CD	ARG	B	143	50.336	-3.548	21.732	1.00	33.24	C
ANISOU	2995	CD	ARG	B	143	4048	3937	4345	22	4	-22 C
ATOM	2996	NE	ARG	B	143	51.169	-3.143	22.929	1.00	39.64	N
ANISOU	2996	NE	ARG	B	143	4844	4732	5465	2	3	-20 N
ATOM	2997	CZ	ARG	B	143	50.982	-3.580	24.193	1.00	49.92	C
ANISOU	2997	CZ	ARG	B	143	6136	6040	6790	-27	-37	-4 C
ATOM	2998	NH1	ARG	B	143	49.275	-4.421	24.429	1.00.	48.96	N
ANISOU	2998	NH1	ARG	B	143	6018	5934	6652	-43	-74	15 N
ATOM	2999	NH2	ARG	B	143	51.755	-3.176	25.229	1.00	54.90	N
ANISOU	2999	NH2	ARG	B	143	6755	6356	7440	-42	-40	-3 N
ATOM	3000	N	TYR	B	144	50.046	-5.485	15.507	1.30	26.13	N
ANISOU	3000	N	TYR	B	144	3198	3057	3374	118	19	-114 N
ATOM	3001	CA	TYR	B	144	49.107	-6.010	15.548	1.00	23.02	C
ANISOU	3001	CA	TYR	B	144	2816	2692	3237	128	-3	-131 C
ATOM	3002	C	TYR	B	144	43.221	-7.551	15.641	1.00	30.09	C
ANISOU	3002	C	TYR	B	144	3711	3551	4170	107	-56	-165 C
ATOM	3003	O	TYR	B	144	48.190	-8.277	15.641	1.00	30.09	O
ANISOU	3003	O	TYR	B	144	3227	3076	3667	87	-100	-174 O
ATOM	3004	CB	TYR	B	144	49.460	-5.549	14.091	1.00	25.70	C
ANISOU	3004	CB	TYR	B	144	3174	3056	3533	172	43	-149 C
ATOM	3005	CG	TYR	B	144	48.453	-6.046	13.084	1.00	24.35	C
ANISOU	3005	CG	TYR	B	144	3016	2927	3309	184	15	-170 C
ATOM	3006	CD1	TYR	B	144	47.093	-5.739	13.254	1.00	27.53	C
ANISOU	3006	CD1	TYR	B	144	3425	3345	3637	74	-9	-145 C

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ATOM	3007	CD2	TYR	B	144	48.851	-6.710	11.900	1.00	27.47	C	
ANISOU	3007	CD2	TYR	B	144	3427	3327	3664	203	17	-218	C
ATOM	3008	CE1	TYR	B	144	46.156	-6.123	12.320	1.00	27.23	C	
ANISOU	3008	CE1	TYR	B	144	3380	3380	3585	184	-38	-165	C
ATOM	3009	CE2	TYR	B	144	47.802	-7.082	10.567	1.00	25.88	C	
ANISOU	3009	CE2	TYR	B	144	3237	3171	3425	217	-11	-241	C
ATOM	3010	CZ	TYR	B	144	46.591	-6.732	11.165	1.00	27.70	C	
ANISOU	3010	CZ	TYR	B	144	3458	3443	3523	205	-39	-214	C
ATOM	3011	OH	TYR	B	144	45.675	-7.236	10.234	1.00	31.27	O	
ANISOU	3011	OH	TYR	B	144	3915	3944	4021	211	-74	-243	O
ATOM	3012	N	LEU	B	145	50.452	-4.066	15.867	1.00	27.03	N	
ANISOU	3012	N	LEU	B	145	3319	3118	3335	112	-54	-185	N
ATOM	3013	CA	LEU	B	145	50.604	-9.548	15.905	1.00	25.71	C	
ANISOU	3013	CA	LEU	B	145	3156	2904	3707	100	-104	-217	C
ATOM	3014	C	LEU	B	145	49.854	-10.147	17.056	1.00	27.37	C	
ANISOU	3014	C	LEU	B	145	3360	3007	3941	53	-155	-187	C
ATOM	3015	O	LEU	B	145	49.120	-11.115	16.946	1.00	28.61	O	
ANISOU	3015	O	LEU	B	145	3527	3241	4104	28	-199	-200	O
ATOM	3016	CB	LEU	B	145	52.137	-9.914	15.908	1.00	26.50	C	
ANISOU	3016	CB	LEU	B	145	3247	2961	3460	125	-37	-243	C
ATOM	3017	CG	LEU	B	145	52.570	-11.403	16.119	1.00	28.06	C	
ANISOU	3017	CG	LEU	B	145	44.50	3796	1115	122	-135	-272	C
ATOM	3018	CD1	LEU	B	145	52.057	-12.247	14.964	1.00	30.69	C	
ANISOU	3018	CD1	LEU	B	145	3807	3425	4428	134	-152	-324	C
ATOM	3019	CD2	LEU	B	145	54.162	-11.410	16.185	1.00	26.65	C	
ANISOU	3019	CD2	LEU	B	145	3250	2850	3945	157	-108	-290	C
ATOM	3020	N	ALA	B	146	50.005	-9.546	16.223	1.00	26.25	N	
ANISOU	3020	N	ALA	B	146	3203	2958	3813	36	-150	-146	N
ATOM	3021	CA	ALA	B	146	49.322	-10.068	19.405	1.00	26.28	C	
ANISOU	3021	CA	ALA	B	146	3265	3015	3397	-9	-194	-110	C
ATOM	3022	C	ALA	B	146	47.776	-10.002	19.265	1.00	27.28	C	
ANISOU	3022	C	ALA	B	146	3327	3123	3916	-36	-209	-96	C
ATOM	3023	O	ALA	B	146	47.050	-10.960	19.630	1.00	26.60	O	
ANISOU	3023	O	ALA	B	146	3240	3022	3843	-75	-253	-68	O
ATOM	3024	CB	ALA	B	146	49.713	-9.274	20.501	1.00	27.69	C	
ANISOU	3024	CB	ALA	B	146	3390	3164	4044	-19	-180	-73	C
ATOM	3025	N	TYR	B	147	47.279	-8.398	18.734	1.00	28.23	N	
ANISOU	3025	N	TYR	B	147	3450	3303	3992	-14	-173	-91	N
ATOM	3026	CA	TYR	B	147	45.823	-8.779	18.376	1.00	27.05	C	
ANISOU	3026	CA	TYR	B	147	3287	3201	3791	-27	-186	-83	C

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ATOM	3027	C	TYR	B	147	45.406	-9.943	17.450	1.00	25.05	C
ANISOU	3027	C	TYR	B	147	3043	2936	3538	-39	-225	-125 C
ATOM	3028	O	TYR	B	147	44.404	-10.631	17.727	1.00	30.01	O
ANISOU	3028	O	TYR	B	147	3661	3571	4170	-83	-267	-119 O
ATOM	3029	CB	TYR	B	147	45.572	-7.406	17.635	1.00	26.90	C
ANISOU	3029	CB	TYR	B	147	3269	2234	3717	16	-138	-78 C
ATOM	3030	CG	TYR	B	147	44.193	-7.406	16.934	1.00	219.58	C
ANISOU	3030	CG	TYR	B	147	3599	3032	4007	15	-157	-80 C
ATOM	3031	CD1	TYR	B	147	43.041	-7.345	17.641	1.00	27.11	C
ANISOU	3031	CD1	TYR	B	147	3261	3364	3676	-5	-167	-46 C
ATOM	3032	CD2	TYR	B	147	44.064	-7.744	15.574	1.00	26.73	C
ANISOU	3032	CD2	TYR	B	147	3251	3290	3515	39	-164	-119 C
ATOM	3033	CE	TYR	B	147	1.41.791	-7.041	17.022	1.00	29.50	C
ANISOU	3033	CE1	TYR	B	147	3545	3727	3935	-4	-186	-49 C
ATOM	3034	CE2	TYR	B	147	42.825	-7.808	14.961	1.00	27.47	C
ANISOU	3034	CE2	TYR	B	147	3332	3443	3664	37	-190	-124 C
ATOM	3035	CZ	TYR	B	147	41.6923	-7.469	15.636	1.00	29.45	C
ANISOU	3035	CZ	TYR	B	147	3551	3736	3932	14	-202	-39 C
ATOM	3036	OH	TYR	B	147	40.510	-7.464	15.043	1.00	31.85	O
ANISOU	3036	OH	TYR	B	147	3834	4105	4161	17	-227	-95 O
ATOM	3037	N	ARG	B	148	45.128	-10.132	16.355	1.00	25.50	N
ANISOU	3037	N	ARG	B	148	2119	2273	3592	-3	-211	-169 N
ATOM	3038	CA	ARG	B	148	45.728	-11.173	15.592	1.00	28.94	C
ANISOU	3038	CA	ARG	B	148	3567	3406	4021	-10	-247	-218 C
ATOM	3039	C	ARG	B	148	45.819	-12.564	15.992	1.00	28.87	C
ANISOU	3039	C	ARG	B	148	2540	3304	4350	-53	-296	-229 C
ATOM	3040	O	ARG	B	148	44.985	-13.422	15.740	1.00	28.83	O
ANISOU	3040	O	ARG	B	148	3561	3319	4073	-89	-338	-250 O
ATOM	3041	CB	ARG	B	148	40.604	-11.164	14.092	1.00	30.11	C
ANISOU	3041	CB	ARG	B	148	3738	3552	4152	41	-219	-270 C
ATOM	3042	CG	ARG	B	148	46.514	-9.929	13.157	1.00	25.46	C
ANISOU	3042	CG	ARG	B	148	3058	35135	3940	88	-170	-264 C
ATOM	3043	CD	ARG	B	148	45.117	-9.849	12.446	1.00	28.78	C
ANISOU	3043	CD	ARG	B	148	3566	3517	3851	83	-197	-270 C
ATOM	3044	NE	ARG	B	148	44.846	-11.012	11.603	1.00	28.49	N
ANISOU	3044	NE	ARG	B	148	3544	3473	3809	71	-240	-333 N
ATOM	3045	CZ	ARG	B	148	45.282	-11.168	10.361	1.30	31.901	C
ANISOU	3045	CZ	ARG	B	148	4306	3526	4210	110	-226	-384 C
ATOM	3046	NH1	ARG	B	148	45.962	-10.164	9.743	1.00	32.75	N
ANISOU	3046	NH1	ARG	B	148	4113	4067	4270	163	-167	-370 N

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ATOM	3047	NH2	ARG	B	148	45.019	-12.288	9.729	1.00	31.96	N
ANISOU	3047	NH2	ARG	B	148	4019	3913	4212	94	-270	-447 N
ATOM	3048	N	LEU	B	149	46.842	-12.820	16.795	1.00	31.02	N
ANISOU	3048	N	LEU	B	149	3840	3545	4400	-61	-223	-214 N
ATOM	3049	CA	LEU	B	149	48.677	-14.052	17.549	1.00	30.83	C
ANISOU	3049	CA	LEU	B	149	3824	3455	4436	-90	-339	-237 C
ATOM	3050	C	LEU	B	149	45.731	-14.156	18.506	1.00	28.20	C
ANISOU	3050	C	LEU	B	149	3474	3139	4130	-149	-067	-158 C
ATOM	3051	O	LEU	B	149	45.167	-15.287	18.087	1.00	31.93	O
ANISOU	3051	O	LEU	B	149	3555	3576	4602	-134	-409	-195 O
ATOM	3052	CB	LEU	B	149	43.232	-14.253	18.233	1.00	26.49	C
ANISOU	3052	CB	LEU	B	149	3278	2848	3940	-88	-333	-195 C
ATOM	3053	CG	LEU	B	149	49.425	-14.297	17.251	1.00	30.29	C
ANISOU	3053	CG	LEU	B	149	3770	3308	4431	-10	-305	-250 C
ATOM	3054	CD1	LEU	B	149	50.760	-14.107	17.916	1.00	29.41	C
ANISOU	3054	CD1	LEU	B	149	3647	3167	4362	18	-288	-234 C
ATOM	3055	CD2	LEU	B	149	49.441	-15.604	16.403	1.00	33.44	C
ANISOU	3055	CD2	LEU	B	149	4195	3653	4859	-6	-338	-311 C
ATOM	3056	N	ILE	B	150	45.350	-13.132	19.232	1.00	29.59	N
ANISOU	3056	N	ILE	B	150	3628	3370	4244	-153	-341	-110 N
ATOM	3057	CA	ILE	B	150	44.220	-13.210	20.102	1.00	28.81	C
ANISOU	3057	CA	ILE	B	150	3509	3301	4136	-206	-360	-65 C
ATOM	3058	C	ILE	B	150	42.915	-13.584	19.335	1.00	29.17	C
ANISOU	3058	C	ILE	B	150	3542	3388	4153	-233	-381	-88 C
ATOM	3059	O	ILE	B	150	42.073	-14.167	19.879	1.00	31.44	O
ANISOU	3059	O	ILE	B	150	3816	3073	4456	-291	-413	-65 O
ATOM	3060	CB	ILE	B	150	44.092	-11.897	20.960	1.00	30.02	C
ANISOU	3060	CB	ILE	B	150	3641	3510	4254	-145	-323	-20 C
ATOM	3061	CG1	ILE	B	150	45.187	-11.899	22.322	1.00	31.77	C
ANISOU	3061	CG1	ILE	B	150	3871	3690	4511	-190	-319	7 C
ATOM	3062	CG2	ILE	B	150	42.700	-11.784	21.627	1.00	29.51	C
ANISOU	3062	CG2	ILE	B	150	3543	3501	4164	-243	-332	19 C
ATOM	3063	CD1	ILE	B	150	45.334	-10.543	22.691	1.00	33.66	C
ANISOU	3063	CD1	ILE	B	150	4095	3976	4716	-170	-278	33 C
ATOM	3064	N	ARG	B	151	42.849	-13.027	18.044	1.00	28.32	N
ANISOU	3064	N	ARG	B	151	3438	3319	4034	-192	-367	-132 N
ATOM	3065	CA	ARG	B	151	41.623	-13.245	17.240	1.00	28.22	C
ANISOU	3065	CA	ARG	B	151	3408	3357	3957	-214	-303	-158 C
ATOM	3066	C	ARG	B	151	41.456	-14.739	16.989	1.00	51.57	C
ANISOU	3066	C	ARG	B	151	3885	3759	4466	-262	-443	-195 C

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ATOM	3067	O	ARG	B	151	40.344	-15.266	17.018	1.00	31.94	O
ANISOU	3067	O	ARG	B	151	3873	3784	4475	-317	-477	-195 O
ATOM	3068	CB	ARG	B	151	41.689	-12.458	15.910	1.00	23.61	C
ANISOU	3068	CB	ARG	B	151	3454	3454	3947	-154	-358	-195 C
ATOM	3069	CG	ARG	B	151	41.450	-10.962	16.071	1.00	28.00	C
ANISOU	3069	CG	ARG	B	151	3369	3449	3820	-114	-323	-154 C
ATOM	3070	CD	ARG	B	151	40.145	-10.557	16.595	1.00	28.57	C
ANISOU	3070	CD	ARG	B	151	3399	3583	3873	-146	-331	-110 C
ATOM	3071	NE	ARG	B	151	38.941	-11.222	16.228	1.00	33.10	N
ANISOU	3071	NE	ARG	B	151	3568	3819	4050	-183	-376	-135 N
ATOM	3072	CZ	ARG	B	151	38.329	-10.878	15.072	1.00	30.08	C
ANISOU	3072	CZ	ARG	B	151	3555	3581	3992	-152	-385	-163 C
ATOM	3073	NH1	ARG	B	151	38.663	-9.701	14.416	1.00	26.69	N
ANISOU	3073	NH1	ARG	B	151	3139	3487	3514	-83	-345	-158 N
ATOM	3074	NH2	ARG	B	151	37.209	1-11.573	14.652	1.00	28.33	N
ANISOU	3074	NH2	ARG	B	151	3306	3695	3762	-145	-432	-190 N
ATOM	3075	N	ILE	B	152	42.582	-15.450	16.832	1.00	31.07	N
ANISOU	3075	N	ILE	B	152	3820	3576	4411	-344	-443	-220 N
ATOM	3076	CA	ILE	B	152	42.487	-16.947	16.508	1.00	30.30	C
ANISOU	3076	CA	ILE	B	152	3745	3402	4360	-258	-497	-265 C
ATOM	3077	C	ILE	B	152	42.596	-17.737	17.924	1.00	31.13	C
ANISOU	3077	C	ILE	B	152	3655	3425	4526	-3138	-319	-212 C
ATOM	3078	O	ILE	B	152	42724	-18.950	17.906	1.00	32.75	O
ANISOU	3078	O	ILE	B	152	4088	3550	4801	-369	-555	-234 O
ATOM	3079	CB	ILE	B	152	43.474	-17.405	15.532	1.00	32.52	C
ANISOU	3079	CB	ILE	B	152	4062	3636	4659	-238	-445	-337 C
ATOM	3080	CG1	ILE	B	152	44.070	-17.173	15.972	1.03	31.31	C
ANISOU	3080	CG1	ILE	B	152	3924	3434	4537	-185	-432	-322 C
ATOM	3081	CG2	ILE	B	152	43.316	-16.678	14.240	1.00	31.37	C
ANISOU	3081	CG2	ILE	B	152	3911	3564	4438	-195	-475	-363 C
ATOM	3082	CD1	ILE	B	152	45.430	-18.095	15.190	1.00	37.48	C
ANISOU	3082	CD1	ILE	B	152	4741	4142	5359	-150	-471	-391 C
ATOM	3083	N	GLY	B	153	42.506	-17.065	19.051	1.00	29.31	N
ANISOU	3083	N	GLY	B	153	6133	3236	4295	-347	-498	-143 N
ATOM	3084	CA	GLY	B	153	42.386	-17.788	20.338	1.00	29.64	C
ANISOU	3084	CA	GLY	B	153	3647	3223	4385	-402	-520	-84 C
ATOM	3085	C	GLY	B	153	43.718	-18.191	20.971	1.30	34.47	C
ANISOU	3085	C	GLY	B	153	4352	3823	5111	-372	-521	-65 C
ATOM	3086	O	GLY	B	153	43.734	-18.920	21.909	1.00	35.53	O
ANISOU	3086	O	GLY	B	153	4371	3780	5160	-411	-545	-19 O

TABLE 3-continued

ATOM	3087	N	LYS	B	154	44.559	-17.059	23.493	1.00	29.87	N	
ANISOU	3087	N	LYS	B	154	3678	3136	4421	-393	-495	-95 N	
ATOM	3088	CA	LYS	B	154	46.106	-17.901	21.164	1.00	34.55	C	
ANISOU	3088	CA	LYS	B	154	4337	3719	5108	-271	-495	-76 C	
ATOM	3089	C	LYS	B	154	45.347	-15.924	22.331	1.00	28.92	C	
ANISOU	3089	C	LYS	B	154	3589	3042	4356	-256	-470	-14 C	
ATOM	3090	O	LYS	B	154	45.833	-15.757	22.291	1.00	31.33	O	
ANISOU	3090	O	LYS	B	154	3871	3429	4605	-251	-437	-5 O	
ATOM	3091	CB	LYS	B	154	47.235	-17.815	20.107	1.00	35.05	C	
ANISOU	3091	CB	LYS	B	154	4401	3743	5167	-202	-475	-142 C	
ATOM	3092	CG	LYS	B	154	47.256	-18.971	19.105	1.00	42.07	C	
ANISOU	3092	CG	LYS	B	154	5320	4571	6093	-200	-504	-209 C	
ATOM	3093	CD	LYS	B	154	47.701	-20.227	19.842	1.00	41.39	C	
ANISOU	3093	CD	LYS	B	154	5250	4352	5085	-217	-544	-186 C	
ATOM	3094	CE	LYS	B	154	48.246	-21	225	18.885	1.00	41.24	C
ANISOU	3094	CE	LYS	B	154	5273	4285	8111	-187	-561	-250 C	
ATOM	3095	NZ	LYS	B	154	48.611	-22.377	19.740	1.00	37.78	N	
ANISOU	3095	NZ	LYS	B	154	4862	3743	5751	-202	-600	-224 N	
ATOM	3096	N	LYS	B	155	47.050	-17.353	23.390	1.00	30.73	N	
ANISOU	3096	N	LYS	B	155	3624	3221	4520	-257	-488	30 N	
ATOM	3097	CA	LYS	B	155	47.573	-16.396	24.355	1.00	31.93	C	
ANISOU	3097	CA	LYS	B	155	3961	3423	4750	-250	-455	72 C	
ATOM	3098	C	LYS	B	155	48.714	-15.574	23.789	1.00	34.94	C	
ANISOU	3098	C	LYS	B	155	4336	3811	5130	-187	-434	30 C	
ATOM	3099	O	LYS	B	155	49.723	-15.297	23.415	1.00	35.10	O	
ANISOU	3099	O	LYS	B	155	4405	3398	5324	-150	-446	1 O	
ATOM	3100	CB	LYS	B	155	47.986	-17.118	25.365	1.00	33.73	C	
ANISOU	3100	CB	LYS	B	155	4200	3604	5013	-259	-499	133 C	
ATOM	3101	CG	LYS	B	155	46.855	-17.852	25.302	1.00	35.22	C	
ANISOU	3101	CG	LYS	B	155	4775	4162	6583	-335	-525	184 C	
ATOM	3102	CD	LYS	B	155	45.741	-15.502	26.592	1.00	42.01	C	
ANISOU	3102	CD	LYS	B	155	5225	4743	5995	-358	-492	203 C	
ATOM	3103	CE	LYS	B	155	44.751	-17.412	27.622	1.00	40.75	C	
ANISOU	3103	CE	LYS	B	155	5055	4590	5833	-439	-511	272 C	
ATOM	3104	NZ	LYS	B	155	44.159	-18.665	27.092	1.00	40.53	N	
ANISOU	3104	NZ	LYS	B	155	5054	4495	5850	-435	-544	243 N	
ATOM	3105	N	ALA	B	156	48.603	-14.373	23.675	1.00	30.42	N	
ANISOU	3105	N	ALA	B	156	3742	3337	408	-171	-393	25 N	
ATOM	3106	CA	ALA	B	156	49.770	-13.533	23.209	1.00	32.01	C	
ANISOU	3106	CA	ALA	B	156	5536	3515	4712	-117	-859	-7 C	

TABLE 3-continued

ATOM	3107	C	ALA	B	156	49.902	-12.373	23.982	1.00	32.90	C	
ANISOU	3107	C	ALA	B	156	4028	5682	4791	-115	-329	20 C	
ATOM	3108	O	ALA	B	156	48.873	-11.582	24.243	1.00	34.72	O	
ANISOU	3108	O	ALA	B	156	4251	3966	4975	-139	-314	41 O	
ATOM	3109	CB	ALA	B	156	49.594	-13.246	21.732	1.00	25.91	C	
ANISOU	3109	CB	ALA	B	156	3294	2691	4041	-89	-328	-61 C	
ATOM	3110	N	ILE	B	157	51.121	-12.031	24.310	1.00	33.84	N	
ANISOU	3110	N	ILE	B	157	3755	3410	4551	-85	-319	15 N	
ATOM	3111	CA	ILE	B	157	51.324	-10.721	24.953	1.00	37.37	C	
ANISOU	3111	CA	ILE	B	157	4525	4248	5310	-85	-288	29 C	
ATOM	3112	C	ILE	B	157	52.372	-9.945	24.274	1.30	34.53	C	
ANISOU	3112	C	ILE	B	157	4192	3930	4998	-45	-247	-8 C	
ATOM	3113	O	ILE	B	157	53.388	-10.500	23.915	1.00	35.42	O	
ANISOU	3113	O	ILE	B	157	4299	4335	5153	-19	-255	-32 O	
ATOM	3114	CB	ILE	B	157	51.651	-10.562	26.415	1.00	38.23	C	
ANISOU	3114	CB	ILE	B	157	4566	4397	5464	-105	-318	71 C	
ATOM	3115	CG	ILE	B	157	52.163	-9.557	26.975	1.00	46.50	C	
ANISOU	3115	CG	ILE	B	157	1	5594	5486	6459	-55	-287	68 C
ATOM	3116	CG2	ILE	B	157	52.782	-11.754	25.587	1.00	40.85	C	
ANISOU	3116	CG2	ILE	B	157	4994	4677	5849	-85	-363	71 C	
ATOM	3117	CD1	ILE	B	157	52.357	-9.770	23.443	1.00	36.50	C	
ANISOU	3117	CD1	ILE	B	157	4422	4223	5213	-119	-323	108 C	
ATOM	3118	N	MET	B	158	52.113	-3.608	24.064	1.00	32.05	N	
ANISOU	3118	N	MET	B	158	3873	3659	4647	-42	-201	-13 N	
ATOM	3119	CA	MET	B	158	53.093	-7.769	23.542	1.00	37.18	C	
ANISOU	3119	CA	MET	B	158	4510	4311	5305	-14	-157	-40 C	
ATOM	3120	C	MET	B	158	53.691	-6.957	24.714	1.00	40.40	C	
ANISOU	3120	C	MET	B	158	4898	4735	5717	-27	-153	-26 C	
ATOM	3121	O	MET	B	158	52.944	-5.325	25.478	1.00	40.54	O	
ANISOU	3121	O	MET	B	158	4920	4784	5700	-49	-1150	-3 O	
ATOM	3122	CB	MET	B	158	52.373	-6.855	22.538	1.00	44.54	C	
ANISOU	3122	CB	MET	B	158	4202	4022	4.941	-2	-110	-49 C	
ATOM	3123	CG	MET	B	158	53.218	-5.785	21.923	1.00	33.44	C	
ANISOU	3123	CG	MET	B	158	4041	3874	4790	23	-53	-70 C	
ATOM	3124	SD	MET	B	158	54.786	-6.398	21.257	1.00	34.89	S	
ANISOU	3124	SD	MET	B	158	4207	4024	5027	52	-44	-107 S	
ATOM	3125	CE	MET	B	158	55.517	-4.319	20.535	1.00	33.44	C	
ANISOU	3125	CE	MET	B	158	4392	4237	5218	61	30	-115 C	
ATOM	3126	N	PHE	B	159	55.013	-5.321	24.812	1.00	32.33	N	
ANISOU	3126	N	PHE	B	159	3856	3700	4739	-12	-150	-45 N	

TABLE 3-continued

ATOM	3127	CA	PHE	B	159	55.724	-6.035	25.302	1.00	37.72	C	
ANISOU	3127	CA	PHE	B	159	4512	4397	5424	-26	-146	-43	C
ATOM	3128	C	PHE	B	159	56.4138	-4.890	25.037	1.00	40.40	C	
ANISOU	3128	C	PHE	B	159	4341	4799	5775	-14	-34	-73	C
ATOM	3129	O	PHE	B	159	57.336	-5.118	24.204	1.00	60.48	O	
ANISOU	3129	O	PHE	B	159	6008	6000	7031	10	-63	-95	O
ATOM	3130	CB	PHE	B	159	56.886	-6.939	26.369	1.00	29.35	C	
ANISOU	3130	CB	PHE	B	159	3425	3315	4412	-16	-190	-45	C
ATOM	3131	CG	PHE	B	159	56.445	-8.147	27.107	1.00	31.23	C	
ANISOU	3131	CG	PHE	B	159	3580	3544	4658	-26	-253	-10	C
ATOM	3132	CD1	PHE	B	159	55.310	-8.065	28.544	1.00	33.02	C	
ANISOU	3132	CD1	PHE	B	159	3900	3789	4358	-52	-283	22	C
ATOM	3133	CD2	PHE	B	159	56.353	-9	376	26.551	1.00	29.20	C
ANISOU	3133	CD2	PHE	B	159	3434	3240	4422	-9	-277	-11	C
ATOM	3134	CE1	PHE	B	159	55.972	-9.206	29.270	1.00	30.93	C	
ANISOU	3134	CE1	PHE	B	159	3545	3508	4595	-61	-843	63	C
ATOM	3135	CE2	PHE	B	159	55.021	-10.526	27.277	1.90	33.02	C	
ANISOU	3135	CE2	PHE	B	159	3680	3445	4652	-20	-334	26	C
ATOM	3136	CZ	PHE	B	159	55.850	-10.422	26.566	1.00	32.56	C	
ANISOU	3136	CZ	PHE	B	159	3372	3666	4834	-46	-366	68	C
ATOM	3137	N	GLU	B	160	55.157	-3.534	25.414	1.00	49.66	N	
ANISOU	3137	N	GLU	B	160	5015	5934	6929	-31	-50	-71	N
ATOM	3138	CA	GLU	B	160	57.253	-2.096	25.161	1.00	46.74	C	
ANISOU	3138	CA	GLU	B	160	5622	5559	6576	-31	-4	-98	C
ATOM	3139	C	GLU	B	160	58.104	-2.277	26.338	1.00	46.17	C	
ANISOU	3139	C	GLU	B	160	5518	5495	6528	-53	-29	-107	C
ATOM	3140	O	GLU	B	160	59.193	-1.526	25.225	1.00	38.33	O	
ANISOU	3140	O	GLU	B	160	4496	4498	5571	-64	4	-134	O
ATOM	3141	CB	GLU	B	160	56.796	-1.250	24.548	1.00	46.74	C	
ANISOU	3141	CB	GLU	B	160	5644	5552	5554	-32	61	-99	C
ATOM	3142	CG	GLU	B	160	55.924	-1.463	23.347	1.00	45.43	C	
ANISOU	3142	CG	GLU	B	160	5508	5397	6356	-5	83	-88	C
ATOM	3143	CD	GLU	B	160	54.503	-1.800	23.766	0.50	43.24	C	
ANISOU	3143	CD	GLU	B	160	5255	5139	6035	-10	52	-63	C
ATOM	3144	OE1	GLU	B	160	54.166	-1.809	24.985	1.00	46.05	O	
ANISOU	3144	OE1	GLU	B	160	5607	5509	6382	-33	19	-52	O
ATOM	3145	OE2	GLU	B	160	53	710	-2.025	22.362	1.00	45.52	O
ANISOU	3145	OE2	GLU	B	160	5714	5436	5347	9	61	-56	O
ATOM	3146	N	ASP	B	161	57.623	-2.763	27.572	1.00	34.01	N	
ANISOU	3146	N	ASP	B	161	3933	3972	4967	-72	-84	-86	N

TABLE 3-continued

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ATOM	3147	CA	ASP	B	161	56.504	-2.565	28.775	1.00	30.20	C
ANISOU	3147	CA	ASP	B	161	3469	3505	4501	-91	-121	-96 C
ATOM	3148	C	ASP	B	161	59.503	-3.321	28.518	1.50	31.12	C
ANISOU	3148	C	ASP	B	161	3555	3508	4664	-68	-160	-97 C
ATOM	3149	O	ASP	B	161	59.128	-4.988	28.870	1.00	30.34	O
ANISOU	3149	O	ASP	B	161	3429	3462	4523	-57	-207	-69 O
ATOM	3150	CB	ASP	B	161	57.750	-2.734	30.114	1.00	27.95	C
ANISOU	3150	CB	ASP	B	161	3203	3255	4174	-113	-163	-71 C
ATOM	3151	CG	ASP	B	161	58.639	2.528	31.317	1.00	30.53	C
ANISOU	3151	CG	ASP	B	161	3493	3632	4534	-132	-202	-86 C
ATOM	3152	OD1	ASP	B	161	56.266	-2.033	32.372	1.00	32.03	O
ANISOU	3152	OD1	ASP	B	161	3694	3824	4653	-154	-215	-84 O
ATOM	3153	OD2	ASP	B	161	59.923	-2.832	31.158	1.00	34.59	O
ANISOU	3153	OD2	ASP	B	161	3980	4122	5880	-122	-217	-104 O
ATOM	3154	N	THR	B	162	60.691	-3.507	28.155	1.00	30.41	N
ANISOU	3154	N	THR	B	162	3423	3511	4619	-61	-136	-129 N
ATOM	3155	CA	THR	B	162	61.745	-4.495	27.905	1.30	32.98	C
ANISOU	3155	CA	THR	B	162	3711	3825	4995	-31	-165	-137 C
ATOM	3156	C	THR	B	162	62.303	-6.070	29.192	1.00	31.52	C
ANISOU	3156	C	THR	B	162	4257	4418	5561	-33	-237	-125 C
ATOM	3157	O	THR	B	162	63.595	-6.364	29.139	1.00	36.56	O
ANISOU	3157	O	THR	B	162	4137	4265	5501	-1	-273	-124 O
ATOM	3158	CB	THR	B	162	62.862	-3.882	25.992	1.00	31.59	C
ANISOU	3158	CB	THR	B	162	3493	3546	4864	-24	-109	-176 C
ATOM	3159	OG1	THR	B	162	55.195	-2.554	27.476	1.00	35.14	O
ANISOU	3159	OG1	THR	B	162	3925	4114	5313	-64	-82	-195 O
ATOM	3160	CG2	THR	B	162	62.316	-3.749	25.501	1.00	36.35	C
ANISOU	3160	CG2	THR	B	162	4130	4231	5455	-5	-45	-179 C
ATOM	3161	N	HIS	B	163	62.074	-4.426	30.341	1.30	29.44	N
ANISOU	3161	N	HIS	B	163	3238	3426	4523	-57	-256	-119 N
ATOM	3162	CA	HIS	B	163	52.413	-5.054	31.534	1.00	31.33	C
ANISOU	3162	CA	HIS	B	163	3420	3652	4719	-67	-833	-99 C
ATOM	3163	C	HIS	B	163	61.542	-5.238	31.556	1.00	32.11	C
ANISOU	3163	C	HIS	B	163	3549	3771	4630	-53	-373	-48 C
ATOM	3164	O	HIS	B	163	61.496	-7.352	32.159	1.00	27.42	O
ANISOU	3164	O	HIS	B	163	2994	3166	4260	-27	-428	-23 O
ATOM	3165	CB	HIS	B	163	62.429	-4.117	32.744	1.00	27.75	C
ANISOU	3165	CB	HIS	B	163	3000	3279	4256	-165	-346	-110 C
ATOM	3166	CG	HIS	B	163	63.557	-3.093	32.712	1.00	30.62	C
ANISOU	3166	CG	HIS	B	163	3311	3657	4655	-125	-322	-152 C

TABLE 3-continued

ATOM	3167	ND1	HIS	B	163	64.391	-2.933	33.770	1.30	33.23	N
ANISOU	3167	ND1	HIS	B	163	3223	3549	4518	-140	-373	-178 N
ATOM	3168	CD2	HIS	B	163	63.985	-2.227	31.744	1.00	27.82	C
ANISOU	3168	CD2	HIS	B	163	2940	8284	4345	-135	-255	-156 C
ATOM	3169	CE1	HIS	B	163	65.284	-1.969	33.515	1.00	31.97	C
ANISOU	3169	CE1	HIS	B	163	3396	5675	4678	-164	-339	-227 C
ATOM	3170	NE2	HIS	B	163	65.039	-1.523	32.252	1.30	26.59	N
ANISOU	3170	NE2	HIS	B	163	2985	3407	4471	-162	-264	-237 N
ATOM	3171	N	LEU	B	164	60.225	-5.974	31.803	1.00	33.33	N
ANISOU	3171	N	LEU	B	164	3422	3541	4556	-73	-949	-29 N
ATOM	3172	CA	LEU	B	164	59.245	-7.078	32.018	1.03	29.71	C
ANISOU	3112	CA	LEU	B	164	3332	3452	4454	-70	-383	21 C
ATOM	3173	C	LEU	B	164	50.429	-8.203	30.974	1.00	31.91	C
ANISOU	3173	C	LEU	B	164	3665	3678	4780	-35	-387	22 C
ATOM	3174	O	LEU	B	164	59.317	-9.373	81.312	1.00	29.96	O
ANISOU	3174	O	LEU	B	164	3432	5437	4345	-24	-435	55 O
ATOM	3175	CB	LEU	B	164	57.826	-6.544	33.829	1.00	29.02	C
ANISOU	3175	CB	LEU	B	164	3335	3375	4316	-34	-344	31 C
ATOM	3176	CG	LEU	B	164	57.155	-5.769	32.909	1.00	36.97	C
ANISOU	3176	CG	LEU	B	164	4354	4429	5263	-126	-343	41 C
ATOM	3177	CD1	LEU	B	164	55.651	-5.963	32.553	1.00	39.34	C
ANISOU	3177	CD1	LEU	B	164	4691	4734	5523	-138	-312	61 C
ATOM	3178	CD2	LEU	B	164	57.277	-6.541	34.216	1.00	33.34	C
ANISOU	3178	CD2	LEU	B	164	3394	3993	4751	-134	-409	83 C
ATOM	3179	N	ALA	B	165	59.699	-7.831	29.720	1.09	23.65	N
ANISOU	3179	N	ALA	B	165	3244	3247	4394	-18	-335	-17 N
ATOM	3180	CA	ALA	B	165	59.915	-9.662	23.683	1.30	32.22	C
ANISOU	3180	CA	ALA	B	165	3702	3654	4883	18	-335	-26 C
ATOM	3181	C	ALA	B	165	61.163	-9.693	28.988	1.00	33.43	C
ANISOU	3181	C	ALA	B	165	3437	3409	4716	53	-380	-28 C
ATOM	3182	O	ALA	B	165	51.173	-10.939	23.797	1.00	28.11	O
ANISOU	3182	O	ALA	B	165	3146	3058	4440	82	-414	-14 O
ATOM	3183	CB	ALA	B	165	60.107	-8.195	27.321	1.00	32.39	C
ANISOU	3183	CB	ALA	B	165	3710	3663	4912	31	-267	-69 C
ATOM	3184	N	ALA	B	166	62.239	-9.029	29.454	1.00	27.28	N
ANISOU	3184	N	ALA	B	166	2955	3043	4334	53	-382	-49 N
ATOM	3185	CA	ALA	B	166	63.436	-9.815	29.812	1.30	30.73	C
ANISOU	3185	CA	ALA	B	166	3381	9472	4823	91	-430	-49 C
ATOM	3186	C	ALA	B	166	63.085	-10.755	30.956	1.00	29.20	C
ANISOU	3186	C	ALA	B	166	3210	3279	4614	92	-504	8 C

TABLE 3-continued

ATOM	3187	O	ALA	B	166	63.445	-11.949	30.946	1.00	27.97	O
ANISOU	3187	O	ALA	B	166	3056	3374	4499	133	-547	27 O
ATOM	3188	CB	ALA	B	166	64.633	-8.333	30.256	1.00	23.92	C
ANISOU	3188	CB	ALA	B	166	3212	3416	4740	85	-428	-80 C
ATOM	3189	N	MET	B	167	62.384	-10.251	31.967	1.03	28.75	N
ANISOU	3189	N	MET	B	167	3175	3250	4499	53	-520	37 N
ATOM	3190	CA	MET	B	167	62.064	-11.154	33.116	1.00	34.35	C
ANISOU	3190	CA	MET	B	167	3140	3137	4426	49	-588	100 C
ATOM	3191	C	MET	B	167	81.173	-12.275	32.677	1.00	29.24	C
ANISOU	3191	C	MET	B	167	2313	3245	4549	53	-600	135 C
ATOM	3192	O	MET	B	167	51.429	-13.492	33.085	1.00	29.78	O
ANISOU	3192	O	MET	B	167	3391	3279	4644	84	-653	177 O
ATOM	3193	CB	MET	B	167	61.313	-10.329	94.117	1.00	27.29	C
ANISOU	3193	CB	MET	B	167	3032	3118	4219	1	-587	119 C
ATOM	3194	CG	MET	B	167	62.330	-3.264	34.730	1.00	29.36	C
ANISOU	3194	CG	MET	B	167	3242	3441	4474	-9	-594	32 C
ATOM	3195	SD	MET	B	167	61.237	-8.486	26.051	1.00	32.25	S
ANISOU	3195	SD	MET	B	167	3642	3857	4745	-52	-598	106 S
ATOM	3196	CE	MET	B	167	42.496	-7.594	37.027	1.00	28.46	C
ANISOU	3196	CE	MET	B	167	3116	3.465	4270	-71	-630	66 C
ATOM	3197	N	SER	B	168	50.155	-12.127	31.817	1.00	27.94	N
ANISOU	3197	N	SER	B	168	3174	3059	4359	36	-550	122 N
ATOM	3198	CA	SER	B	168	59.356	-13.261	31.273	1.30	29.13	C
ANISOU	3198	CA	SER	B	168	3378	3156	4524	39	-559	145 C
ATOM	3199	C	SER	B	168	60.177	-24.180	30.445	1.30	29.11	C
ANISOU	3199	C	SER	B	168	3365	3093	4601	91	-570	119 C
ATOM	3200	O	SER	B	168	20.065	-15.409	30.626	1.00	33.58	O
ANISOU	3200	O	SER	B	168	3591	3235	44320	107	-614	154 O
ATOM	3201	CB	SER	B	168	58.207	-12.755	30.377	1.00	32.24	C
ANISOU	3201	CB	SER	B	168	3799	3554	4336	10	-505	123 C
ATOM	3202	OG	SER	B	168	57.114	-12.606	31.150	1.00	47.36	O
ANISOU	3202	OG	SER	B	168	5740	5497	6759	-34	-512	166 O
ATOM	3203	N	ALA	B	169	91.019	-13.620	29.537	1.60	29.49	N
ANISOU	3203	N	ALA	B	169	3369	3144	4670	120	-523	59 N
ATOM	3204	CA	ALA	B	169	61.853	-14.441	28.651	1.00	31.63	C
ANISOU	3204	CA	ALA	B	169	2632	3372	5014	175	-529	25 C
ATOM	3205	C	ALA	B	169	62.755	-15.304	29.487	1.00	31.88	C
ANISOU	3205	C	ALA	B	169	3642	3382	5089	218	-594	56 C
ATOM	3206	O	ALA	B	169	63.036	-16.481	29.088	1.00	30.91	O
ANISOU	3206	O	ALA	B	169	3539	3191	5019	261	-619	54 O

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ATOM	3207	CB	ALA	B	169	62.713	-13.641	27.638	1.00	30.32	C
ANISOU	3207	CB	ALA	B	169	3421	3233	4357	199	-471	-39 C
ATOM	3208	N	SER	B	170	53.213	-14.793	30.652	1.00	32.38	N
ANISOU	3208	N	SER	B	170	3674	3498	5130	204	-625	83 N
ATOM	3209	CA	SER	B	170	64.189	-15.524	31.444	1.00	34.59	C
ANISOU	3209	CA	SER	B	170	3925	3770	5448	248	-590	113 C
ATOM	3210	C	SER	B	170	63.599	-16.740	32.162	1.00	35.48	C
ANISOU	3210	C	SER	B	170	4217	3955	5688	250	-750	185 C
ATOM	3211	O	SER	B	170	64.378	-17.555	32.594	1.00	32.72	O
ANISOU	3211	O	SER	B	170	3727	3454	5253	300	-805	213 O
ATOM	3212	CB	SER	B	170	64.887	-14.609	32.474	1.00	31.02	C
ANISOU	3212	CB	SER	B	170	3421	3398	4968	233	-712	118 C
ATOM	3213	OG	SER	B	170	64.341	-14.452	33.522	1.00	29.57	O
ANISOU	3213	OG	SER	B	170	3274	3248	4718	187	-742	173 O
ATOM	3214	N	ARG	B	171	52.257	-16.816	322.318	1.00	33.71	N
ANISOU	3214	N	ARG	B	171	3925	3593	5291	1197	-741	220 N
ATOM	3215	CA	ARG	B	171	61.603	-17.997	32.983	1.00	32.29	C
ANISOU	3215	CA	ARG	B	171	3798	3356	5110	189	-792	295 C
ATOM	3216	C	ARG	B	171	50.813	-18.803	31.952	1.00	33.18	C
ANISOU	3216	C	ARG	B	171	3960	3988	5258	183	-771	280 C
ATOM	3217	O	ARG	B	171	60.023	-19.885	32.269	1.90	34.09	O
ANISOU	3217	O	ARG	B	171	4126	3450	5376	159	-797	333 O
ATOM	3218	CB	ARG	B	171	60.761	-17.580	84.168	1.00	33.47	C
ANISOU	3218	CB	ARG	B	171	3971	3551	5180	131	-806	358 C
ATOM	3219	CG	ARG	B	171	61.548	-18.659	35.081	1.00	35.24	C
ANISOU	3219	CG	ARG	B	171	4272	3998	5499	134	-823	353 C
ATOM	3220	CD	ARG	B	171	52.827	-17.396	35.500	1.00	40.88	C
ANISOU	3220	CD	ARG	B	171	4828	4367	6139	199	-885	872 C
ATOM	3221	NE	ARG	B	171	62.529	-18.533	36.350	1.00	47.43	N
ANISOU	3221	NE	ARG	B	171	5702	5354	5967	206	-944	460 N
ATOM	3222	CZ	ARG	B	171	53.262	-19.642	36.430	1.00	57.64	C
ANISOU	3222	CZ	ARG	B	171	6996	5584	8322	268	-995	491 C
ATOM	3223	NH1	ARG	B	171	64.317	-19.840	35.622	1.00	54.44	N
ANISOU	3223	NH1	ARG	B	171	6540	6149	7988	332	-952	434 N
ATOM	3224	NH2	ARG	B	171	52.895	-20.677	37.291	1.00	57.55	N
ANISOU	3224	NH2	ARG	B	171	7043	6548	8312	267	-1046	582 N
ATOM	3225	N	SER	B	172	51.100	-18.550	30.666	1.00	31.33	N
ANISOU	3225	N	SER	B	172	3679	3110	5021	205	-724	205 N
ATOM	3226	CA	SER	B	172	60.315	-19.148	29.647	1.00	33.19	C
ANISOU	3226	CA	SER	B	172	3937	3315	6307	195	-701	177 C

TABLE 3-continued

ATOM	3227	C	SER	B	172	60.629	-20.587	29.595	1.00	35.80	C
ANISOU	3227	C	SER	B	172	4477	3608	5837	240	-749	197 C
ATOM	3228	O	SER	B	172	51.787	-21.137	25.900	1.00	31.56	O
ANISOU	3228	O	SER	B	172	3735	2984	5223	304	-784	203 O
ATOM	3229	CB	SER	B	172	60.700	-18.410	28.393	1.00	34.41	C
ANISOU	3229	CB	SER	B	172	4114	3495	5467	213	-641	95 C
ATOM	3230	OG	SER	B	172	59.902	-18.718	27.289	1.00	38.83	O
ANISOU	3230	OG	SER	B	172	4709	4019	6026	204	-610	59 O
ATOM	3231	N	SER	B	173	5.625	-21.461	29.153	1.00	43.24	N
ANISOU	3231	N	SER	B	173	4967	4037	6284	210	-751	202 N
ATOM	3232	CA	SER	B	173	59.591	-22.935	29.252	1.03	37.56	C
ANISOU	3232	CA	SER	B	173	4584	3000	6024	235	-799	223 C
ATOM	3233	C	SER	B	173	59.989	-23.577	28.051	1.00	37.29	C
ANISOU	3233	C	SER	B	173	4680	3483	6007	222	-778	177 C
ATOM	3234	O	SER	B	173	58.348	-22.892	27.250	1.00	33.92	O
ANISOU	3234	O	SER	B	173	4249	3100	5538	139	-731	128 O
ATOM	3235	CB	SER	B	173	59.070	-23.392	30.55	1.00	36.29	C
ANISOU	3235	CB	SER	B	173	4545	3414	5830	190	-845	335 C
ATOM	3236	OG	SER	B	173	57.74	-23.091	30.496	1.00	35.58	O
ANISOU	3236	OG	SER	B	173	4431	3348	5639	111	-819	347 O
ATOM	3237	N	GLN	B	174	59.206	-24.887	27.863	1.00	35.72	N
ANISOU	3237	N	GLN	B	174	4519	3172	5881	257	-813	179 N
ATOM	3238	CA	GLN	B	174	58.627	-25.598	26.725	1.00	95.52	C
ANISOU	3238	CA	GLN	B	174	4535	3071	5890	247	-800	117 C
ATOM	3239	C	GLN	B	174	57.125	-25.253	261.605	1.00	34.43	C
ANISOU	3239	C	GLN	B	174	4422	2964	5694	153	-780	128 C
ATOM	3240	O	GLN	B	174	56.395	-25.212	27.592	1.00	56.42	O
ANISOU	3240	O	GLN	B	174	4689	3235	5315	93	-799	207 O
ATOM	3241	CB	GLN	B	174	58.917	-27.187	26.801	1.00	37.78	C
ANISOU	3241	CB	GLN	B	174	4873	3213	6273	286	-850	134 C
ATOM	3242	CG	GLN	B	174	50.402	-27.466	26.625	1.00	51.55	C
ANISOU	3242	CG	GLN	B	174	6583	4931	6071	352	-860	101 C
ATOM	3243	CD	GLN	B	174	53.888	-26.979	25.258	1.00	51.17	C
ANISOU	3243	CD	GLN	B	174	6502	4920	5020	435	-803	-13 C
ATOM	3244	OE1	GLN	B	174	60.301	-27.351	24.233	1.00	60.08	O
ANISOU	3244	OE1	GLN	B	174	7655	6005	9153	418	-785	-77 O
ATOM	3245	NE2	GLN	B	174	61.867	-26.095	25.219	1.00	51.51	N
ANISOU	3245	NE2	GLN	B	174	6478	5049	5045	481	-779	-37 N
ATOM	3246	N	GLY	B	175	56.707	-24.936	25.395	1.00	32.48	N
ANISOU	3246	N	GLY	B	175	4173	2755	5426	143	-740	47 N

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ATOM	3247	CA	GLY	B	175	55.374	-24.467	25.138	1.00	34.92	C	
ANISOU	3247	CA	GLY	B	175	4435	3092	5580	65	-714	45 C	
ATOM	3248	C	GLY	B	175	65.327	-22.995	24.752	1.00	34.94	C	
ANISOU	3248	C	GLY	B	175	4453	3216	5608	62	-666	15 C	
ATOM	3249	O	GLY	B	175	54.314	-22.504	24.256	1.00	52.45	O	
ANISOU	3249	O	GLY	B	175	4140	2947	5244	14	-641	-5 O	
ATOM	3250	N	ASP	B	176	56.310	-22.248	25.134	1.00	33.52	N	
ANISOU	3250	N	ASP	B	176	4230	3088	5419	108	-652	20 N	
ATOM	3251	CA	ASP	B	176	56.422	-20.793	24.848	1.00	33.17	C	
ANISOU	3251	CA	ASP	B	176	4144	3149	5311	105	-800	-5 C	
ATOM	3252	C	ASP	B	176	67.051	-20.582	23.495	1.00	36.77	C	
ANISOU	3252	C	ASP	B	176	4585	3610	5775	156	-559	-92 C	
ATOM	3253	O	ASP	B	176	57.821	-21.462	23.003	1.00	32.82	O	
ANISOU	3253	O	ASP	B	176	4219	3169	5461	211	-572	-132 O	
ATOM	3254	CB	ASP	B	176	57.283	-23.076	25.862	1.00	33.56	C	
ANISOU	3254	CB	ASP	B	176	4165	3265	5361	124	-604	35 C	
ATOM	3255	CG	ASP	B	176	56.711	-20.100	27.260	1.00	34.30	C	
ANISOU	3255	CG	ASP	B	176	4257	3331	5416	75	-339	122 C	
ATOM	3256	OD	ASP	B	176	1	55.635	-20.485	27.424	1.00	24.18	O
ANISOU	3256	OD1	ASP	B	176	4275	3329	5363	13	-649	164 O	
ATOM	3257	OD2	ASP	B	176	57.472	-19.779	28.211	1.00	34.07	O	
ANISOU	3257	OD2	ASP	B	176	4200	3363	5383	92	-458	159 O	
ATOM	3258	N	LEU	B	177	56.391	-19.492	22.835	1.00	31.43	N	
ANISOU	3258	N	LEU	B	177	3892	3009	5040	142	-503	-123 N	
ATOM	3259	CA	LEU	B	177	57.352	-19.045	21.645	1.00	32.52	C	
ANISOU	3259	CA	LEU	B	177	4011	3173	6174	189	-461	-195 C	
ATOM	3260	C	LEU	B	177	57.459	-17.509	21.575	1.00	33.77	C	
ANISOU	3260	C	LEU	B	177	4132	3425	5274	176	-411	-188 C	
ATOM	3261	O	LEU	B	177	56.519	-15.851	22.133	1.00	31.62	O	
ANISOU	3261	O	LEU	B	177	3891	7222	4979	127	-405	-148 O	
ATOM	3262	CB	LEU	B	177	56.544	-15.495	20.413	1.00	33.33	C	
ANISOU	3262	CB	LEU	B	177	4150	3255	520	183	-450	-252 C	
ATOM	3263	CG	LEU	B	177	56.68	-18.362	19.062	1.00	39.50	C	
ANISOU	3263	CG	LEU	B	177	4935	4998	6914	207	-394	-320 C	
ATOM	3264	CD1	LEU	B	177	57.913	-19.385	18.343	1.00	38.31	C	
ANISOU	3264	CD1	LEU	B	177	4750	3900	5896	279	-378	-381 C	
ATOM	3265	CD2	LEU	B	177	55.425	-19.198	18.243	1.00	40.50	C	
ANISOU	3265	CD2	LEU	B	177	5087	4208	5092	171	-400	-351 C	
ATOM	3266	N	TRP	B	178	53.548	-16.957	21.130	1.00	29.32	N	
ANISOU	3266	N	TRP	B	178	3532	2889	4718	225	-365	-227 N	

TABLE 3-continued

ATOM	3267	CA	TRP	B	178	53.774	-15.457	21.179	1.09	35.39	C
ANISOU	3267	CA	TRP	B	178	3607	3575	4734	213	-319	-220 C
ATOM	2268	C	TRP	B	178	58.885	-14.910	19.315	1.00	35.56	C
ANISOU	3268	C	TRP	B	178	4285	3792	5434	235	-258	-274 C
ATOM	3269	O	TRP	B	178	59.499	-15.553	18.919	1.00	35.39	O
ANISOU	3269	O	TRP	B	178	4254	3743	5443	282	-246	-328 O
ATOM	3270	CB	TRP	B	178	59.977	-15.107	21.979	1.00	34.65	C
ANISOU	3270	CB	TRP	B	178	4324	3660	5382	234	-323	-204 C
ATOM	3271	CG	TRP	B	178	59.754	-15.302	23.413	1.00	31.71	C
ANISOU	3271	CG	TRP	B	178	3755	3275	5018	205	-380	-142 C
ATOM	3272	CD1	TRP	B	178	59.884	-16.535	24.055	1.00	33.72	C
ANISOU	3272	CD1	TRP	B	178	4026	3457	5319	220	-441	-114 C
ATOM	3273	CD2	TRP	B	178	59.259	-14.455	24.435	1.00	33.39	C
ANISOU	3273	CD2	TRP	B	178	3902	3540	5186	153	-363	-94 C
ATOM	3274	NE1	TRP	B	178	59.525	-36.519	25.377	1.00	35.14	N
AAISOU	3274	NE1	TRP	B	178	4210	3552	5461	163	-479	-49 N
ATOM	3275	CE2	TRP	B	178	59.152	-15.234	25.664	1.00	33.71	C
ANISOU	3275	CE2	TRP	B	178	4013	3551	5243	145	-445	-39 C
ATOM	3276	CE3	TRP	B	178	59.001	-13.032	24.460	1.00	32.74	C
ANISOU	3276	CE3	TRP	B	178	3555	3520	5054	130	-338	-94 C
ATOM	3277	CZ2	TRP	B	178	56.760	-14.654	26.855	1.00	32.88	C
ANISOU	3277	CZ2	TRP	B	178	3505	2489	5059	135	-453	11 C
ATOM	3278	CZ3	TRP	B	178	50.576	-12.521	25.673	1.00	34.53	C
ANISOU	3278	CZ3	TRP	B	178	4097	3788	5255	911	-357	-48 C
ATOM	3279	CH2	TRP	B	178	53.489	-13.294	26.830	1.00	31.25	C
ANISOU	3279	CH2	TRP	B	178	3635	3347	4845	79	-418	1 C
ATOM	3280	N	PHE	B	179	58.296	-13.735	19.611	1.00	29.95	N
ANISOU	3280	N	PHE	B	179	9575	3141	4566	206	-217	-252 N
ATOM	3281	CA	PHE	B	179	53.606	-12.949	38.433	1.00	31.45	C
ANISOU	3281	CA	PHE	B	179	3757	3372	4324	220	-151	-201 C
ATOM	3282	C	PHE	B	179	59.388	-11.697	18.822	1.05	30.06	C
ANISOU	3282	C	PHE	B	179	3537	3233	4645	225	-159	-233 C
ATOM	3283	O	PHE	B	179	53.953	-10.953	19.767	1.00	28.93	O
ANISOU	3283	O	PHE	B	179	3392	3117	4483	165	-119	-240 O
ATOM	3284	CB	PHE	B	179	57.300	-12.474	17.584	1.00	-31.13	C
ANISOU	3284	CB	PHE	B	179	3751	3354	4713	205	-122	-301 C
ATOM	3285	CG	PHE	B	179	56.639	-13.565	16.9.55	1.50	31.70	C
ANISOU	3285	CG	PHE	B	179	3859	3403	4781	211	-150	-336 C
ATOM	2286	CD1	PHE	B	179	55.791	-14.451	17.629	1.00	29.14	C
ANISOU	3286	CD1	PHE	B	179	3559	3039	4474	177	-221	-333 C

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ATOM	3287	CD2	PHE	B	179	56.376	-13.742	15.597	1.00	31.94	C	
ANISOU	3287	CD2	PHE	B	179	3901	3444	4791	249	-126	-394	C
ATOM	3288	CE1	PHE	B	179	55.219	-15.500	16.973	1.00	31.85	C	
ANISOU	3288	CE1	PHE	B	179	3934	3344	4822	175	-250	-351	C
ATOM	3289	CE2	PHE	B	179	56.258	-14.785	14.907	1.00	33.92	C	
ANISOU	3289	CE2	PHE	B	179	4166	3663	5038	252	-156	-436	C
ATOM	3290	CZ	PHE	B	179	55.449	-15.683	15.500	1.00	32.96	C	
ANISOU	3290	CZ	PHE	B	179	4059	3496	4945	215	-223	-417	C
ATOM	3291	N	ALA	B	180	80.525	-11.476	18.142	1.00	26.84	N	
ANISOU	3291	N	ALA	B	180	3072	2813	4233	262	-63	-319	N
ATOM	3292	CA	ALA	B	180	81.273	-10.242	18.328	1.00	25.59	C	
ANISOU	3292	CA	ALA	B	180	3025	2851	4226	252	-15	-309	C
ATOM	3293	C	ALA	B	180	61.315	-9.361	17.131	1.00	29.71	C	
ANISOU	3293	C	ALA	B	180	3435	3234	4569	255	56	-322	C
ATOM	3294	O	ALA	B	180	81.252	-9.725	18.028	1.00	29.10	O	
ANISOU	3294	O	ALA	B	180	3351	3211	4486	291	88	-362	O
ATOM	3295	CB	ALA	B	180	52.716	-10.616	13.404	1.00	27.10	C	
ANISOU	3295	CB	ALA	B	180	3035	2909	4353	2815	-11	-335	C
ATOM	3296	N	VAL	B	181	60.329	-8.243	17.326	1.00	28.44	N	
ANISOU	3296	N	VAL	B	181	3290	3151	4386	221	79	-259	N
ATOM	3297	CA	VAL	B	181	59.905	-7.459	16.221	1.00	27.55	C	
ANISOU	3297	CA	VAL	B	181	3201	3070	4198	226	138	-291	C
ATOM	3298	C	VAL	B	181	80.872	-6.237	16.251	1.00	29.31	C	
ANISOU	3298	C	VAL	B	181	3450	3377	4499	215	201	-252	C
ATOM	3299	O	VAL	B	181	60.902	-5.461	17.241	1.00	29.89	O	
ANISOU	3299	O	VAL	B	181	3446	3387	4523	130	194	-254	O
ATOM	3300	CB	VAL	B	181	58.448	-6.937	16.413	1.00	28.33	C	
ANISOU	3300	CB	VAL	B	181	3345	3181	4248	198	124	-258	C
ATOM	3301	CG1	VAL	B	181	55.075	-6.040	15.247	1.00	29.10	C	
ANISOU	3301	CG1	VAL	B	181	3462	3315	4280	210	137	-252	C
ATOM	3302	CG2	VAL	B	181	67.458	-8.153	16.553	1.06	30.17	C	
ANISOU	3302	CG2	VAL	B	181	3604	3394	4467	195	53	-261	C
ATOM	3303	N	SER	B	182	81.702	-5.127	15.213	1.00	29.24	N	
ANISOU	3303	N	SER	B	182	3373	2334	4441	243	260	-309	N
ATOM	3304	CA	SER	B	182	82.784	-5.057	15.222	1.00	29.21	C	
ANISOU	3304	CA	SER	B	182	3312	3336	4452	228	323	-303	C
ATOM	3305	C	SER	B	182	63.204	-4.947	13.775	1.00	30.82	C	
ANISOU	3305	C	SER	B	182	3518	3567	4528	260	398	-324	C
ATOM	3306	O	SER	B	182	63.704	-5.912	13.210	1.00	30.20	O	
ANISOU	3306	O	SER	B	182	2425	3489	4560	1300	394	-355	O

TABLE 3-continued

ATOM	3307	CB	SER	B	182	63.988	-5.529	16.336	1.30	26.11	C
ANISOU	3307	CB	SER	B	182	2356	233.1	4132	230	295	-322 C
ATOM	3308	OG	SER	B	182	65.011	-4.522	15.942	1.00	30.52	O
ANISOU	3308	OG	SER	B	182	3363	2.511	4718	210	359	-322 O
ATOM	3309	N	SER	B	183	63.053	-3.765	13.183	1.00	8.03	N
ANISOU	3309	N	SER	B	183	3181	3230	4230	2461	468	-297 N
ATOM	3310	CA	SER	B	183	63.484	-3	596	11.705	1.90	31.72 C
ANISOU	3310	CA	SER	B	183	3854	13735	4662	275	539	-310 C
ATOM	3311	C	SER	B	183	54.955	-3.773	11.860	1.00	34.85	C
ANISOU	3311	C	SER	B	183	3982	4142	5115	2135	578	-340 C
ATOM	3312	O	SER	B	183	65.450	-4.340	10.663	1.00	33.91	O
ANISOU	3312	O	SER	B	183	31855	4048	4982	3213.	614	-375 O
ATOM	3213	CB	SER	B	183	63.133	-2.191	11.231	1.00	31.42	C
ANISOU	3313	CB	SER	B	183	3645	3712	4580	254	606	-262 C
ATOM	3314	OG	SER	B	183	61.760	-2.122	10.900	1.00	31.36	O
ANISOU	3314	OG	SER	B	183	3701	3712	4503	266	564	-241 O
ATOM	3315	N	SER	B	184	65.670	-3.286	12.651	1.00	130.50	N
ANISOU	3315	N	SER	B	184	3394	3590	4641	248	670	-3231 N
ATOM	3316	CA	SER	B	184	67.130	-3.184	12.589	1.00	33	.59 C
ANISOU	3316	CA	SER	B	184	3695	3909	5080	248	5.113	-854 C
ATOM	3317	C	SER	B	184	67.854	-4.417	13.167	1.00	23.05	C
ANISOU	3317	C	SER	B	184	3645	3979	5139	279	557	-395 C
ATOM	3318	O	SER	B	184	69.047	-4.503	12.979	1.00	32.5.6	O
ANISOU	3918	O	SER	B	184	3446	3368	5054	235.590	-423	O
ATOM	3319	CB	SER	B	184	67.741	-2.017	13.435	1.00	29	C
ANISOU	3319	CB	SER	B	184	3079	13407	4551	137	638	-328 C
ATOM	3320	OG	SER	B	184	67.460	-2.239	14.823	1.00	32.15	O
ANISOU	3320	OG	SER	B	184	3466	3773	4978	165	553	-325 O
ATOM	3321	N	GLY	B	185	67.103	-5.154	13.996	1.00	31.75	N
ANISOU	3321	N	GLY	B	185	3441	3712	4910	283	469	-393 N
ATOM	3322	CA	GLY	B	185	67.568	-6.210	14.835	1.00	34.23	C
ANISOU	3322	CA	GLY	B	185	3714	4007	5286	306	400	-417 C
ATOM	3323	C	GLY	B	185	68.848	-5.714	15.651	1.00	35.28	C
ANISOU	3323	C	GLY	B	185	3766	4154	5486	280	402	-417 C
ATOM	3324	O	GLY	B	185	69.727	-6.512	15.980	1.00	33.50	O
ANISOU	3324	O	GLY	B	185	3483	3929	5315	313	372	-445 O
ATOM	3325	N	SER	B	186	68.848	-4.403	16.047	1.00	34.86	N
ANISOU	3325	N	SER	B	186	3706	4110	5429	221	434	-388 N
ATOM	3326	CA	SER	B	186	70.008	-3.781	16.661	1.00	34.20	C
ANISOU	3326	CA	SER	B	186	3542	4047	5407	187	448	-394 C

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ATOM	3327	C	SER	B	186	69.617	-3.000	17.907	1.00	36	63	C
ANISOU	3327	C	SER	B	186	3859	4340	5719	129	405	-368	C
ATOM	3328	O	SER	B	186	70.433	-2.291	18.455	1	00	35.18	O
ANISOU	3328	O	SER	B	186	3617	4170	5578	89	416	-373	O
ATOM	3329	CB	SER	B	186	70.698	-2.859	15.631	1.00	34.57	C	
ANISOU	3329	CB	SER	B	186	3560	4124	5452	170	555	-396	O
ATOM	3330	OG	SER	B	186	71.207	-3.627	14.537	1.00	33.59	O	
ANISOU	3330	OG	SER	B	186	3415	4023	5324	227	596	-428	O
ATOM	3331	N	THR	B	187	68.365	-3.133	18.354	1.00	30.87	N	
ANISOU	3331	N	THR	B	187	3201	3585	4945	124	356	-343	N
ATOM	3332	CA	THR	B	187	67.881	-2.395	19.522	1.00	32.86	C	
ANISOU	3332	CA	THR	B	187	3468	3825	5191	74	319	-320	C
ATOM	3333	C	THR	B	187	68.254	-3.103	20.778	1.00	31.92	O	
ANISOU	3333	C	THR	B	187	3311	3708	5108	77	232	-328	C
ATOM	3334	O	THR	B	187	67.841	-4.255	20.991	1.00	32.42	O	
ANISOU	3334	O	THR	B	187	3397	3757	5164	115	173	-324	O
ATOM	3335	CB	THR	B	187	66.333	-2.210	19.436	1.00	31.57	C	
ANISOU	3335	CB	THR	B	187	3393	3642	4960	70	309	-289	C
ATOM	3336	OG1	THR	B	187	66.115	-1.342	18.345	1.00	40.69	O	
ANISOU	3336	OG1	THR	B	187	4576	4800	6084	64	391	-276	O
ATOM	3337	OG2	THR	B	187	65.876	-1.421	20.599	1.00	35.87	C	
ANISOU	3337	OG2	THR	B	187	3952	4179	5499	23	279	-272	C
ATOM	3338	N	LYS	B	188	69.011	-2.451	21.645	1.00	32.79	N	
ANISOU	3338	N	LYS	B	188	3367	3834	5256	36	220	-336	N
ATOM	3339	CA	LYS	B	188	69.727	-3.188	22.724	1.00	34.61	C	
ANISOU	3339	CA	LYS	B	188	3542	4081	5529	49	141	-348	C
ATOM	3340	C	LYS	B	188	68.794	-3.999	23.582	1.00	34.72	C	
ANISOU	3340	C	LYS	B	188	3605	4077	5510	62	53	-321	C
ATOM	3341	O	LYS	B	188	69.009	-5.194	23.860	1.00	30.97	O	
ANISOU	3341	O	LYS	B	188	3118	3595	5055	108	1	-321	O
ATOM	3342	CB	LYS	B	188	70.633	-2.249	23.492	1.00	36.24	C	
ANISOU	3342	CB	LYS	B	188	3682	4313	5773	-4	141	-366	C
ATOM	3343	CG	LYS	B	188	71.265	-2.833	24.753	1.00	45.40	C	
ANISOU	3343	CG	LYS	B	188	4788	5498	6965	2	50	-373	C
ATOM	3344	CD	LYS	B	188	72.439	-1.984	25.246	1.00	51.54	C	
ANISOU	3344	CD	LYS	B	188	5479	6312	7792	-45	58	-404	C
ATOM	3345	CE	LYS	B	188	72.601	-1.931	26.801	1.00	64.31	C	
ANISOU	3345	CE	LYS	B	188	7072	7955	9408	-71	-32	-407	C
ATOM	3346	NZ	LYS	B	188	72.064	-3.011	27.739	1.00	51.79	N	
ANISOU	3346	NZ	LYS	B	188	5517	6368	7791	-32	-127	-378	N

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ATOM	3347	N	GLU	B	189	67.785	-3.345	24.309	1.00	30.97	N		
ANISOU	3347	N	GLU	B	189	3184	3594	4989	23	41	-298	N	
ATOM	3348	CA	GLU	B	189	56.936	-4.085	25.191	1.00	32.27	C		
ANISOU	3348	CA	GLU	B	189	3390	3748	5122	32	-33	-271	C	
ATOM	3349	C	GLU	B	189	66.018	-5.092	24.501	1	00	29.50	C	
ANISOU	3349	C	GLU	B	189	3096	2.387	4744	71	-40	-253	C	
ATOM	3350	O	GLU	B	189	65.528	-5.382	25.128	1.03	31.93	O		
ANISOU	3350	O	GLU	B	189	3423	3662	5046	69	-105	-232	O	
ATOM	3351	CB	GLU	B	189	66.189	-3.142	26.132	1.00	28.34	C		
ANISOU	3351	CB	GLU	B	189	2926	32.59	4584	-19	-44	-25	C	
ATOM	3352	CG	GLU	B	189	67.268	-2.359	26.954	1.00	33.12	C		
ANISOU	3352	CG	GLU	B	189	3466	3894	5224	-57	-55	-267	C	
ATOM	3353	CD	GLU	B	189	56.708	-1.615	23.129	-1.09	130.29	C		
ANISOU	3353	CD	GLU	B	189	3.132	3547	4828	-101	-84	-273	O	
ATOM	3354	OE1	GLU	B	189	67.491	-1.022	28.903	1.00	36.08	C		
ANISOU	3354	OE1	GLU	B	189	3817	4337	5583	-135	-103	-304	O	
ATOM	3355	OE2	GLU	B	189	65.490	-1.624	28.333	1.09	35.49	O		
ANISOU	3355	OE2	GLU	B	189	3855	4194	5474	-102	-93	-252	O	
ATOM	3356	N	VAL	B	190	65.723	-4.912	23.219	1.33	33.62	N		
ANISOU	3356	N	VAL	B	190	3288	3525	4896	85	25	-262	N	
ATOM	3357	CA	VAL	B	190	64.895	-5.9.25	22.541	1.30	95.59	C		
ANISOU	3357	CA	VAL	B	190	3170	3339	4711	121	12	-255	C	
ATOM	3358	C	VAL	B	190	95.752	-7.191	22.340	1.00	29.54	C		
ANISOU	3358	C	VAL	B	190	3142	3323	4753	172	-19	-276	C	
ATOM	3359	O	VAL	B	190	65.322	-8.309	22.554	1.00	29.81	O		
ANISOU	3359	O	VAL	B	190	3233	3328	4795	197	-72	-255	O	
ATOM	3360	CB	VAL	B	190	64.345	-5.372	21.227	1.00	31.47	C		
ANISOU	3360	CB	VAL	B	190	3439	3533	4916	124	35	-260	C	
ATOM	3361	CG1	VAL	B	190	53.583	-0.439	20.452	1.00	27.52	C		
ANISOU	2361	CG1	VAL	B	190	3004	13061	4392	155	71	-204	C	
ATOM	3362	CG2	VAL	B	190	63.384	-4.183	21.482	1.00	28.32	C		
ANISOU	3362	CG2	VAL	B	190	3139	3190	4469	82	1.09	-235	C	
ATOM	3363	N	ILE	B	191	66.967	43.029	21.884	1.00	31.55	N		
ANISOU	3363	N	ILE	B	191	3248	3608	5069	190	18	-1306	N	
ATOM	3364	CA	ILE	B	191	67.651	-8.140	21.602	1.00	32.64	C		
ANISOU	3364	CA	ILE	B	191	3434	3724	5245	247	-3	-732	C	
ATOM	3365	C	ILE	B	191	68.133	-9.386	22.204	1.00	33.10	C		
ANISOU	3365	C	ILE	B	191	3405	377	2	5334	257	-52	-312	C
ATOM	3366	O	ILE	B	191	68.275	-10.160	22.928	1.00	3481	O		
ANISOU	3366	O	ILE	B	191	3691	3950	5550	308	-178	-314	O	

TABLE 3-continued

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ATOM	3367	CB	ILE	B	191	69.237	-7.613	21.150	1.00	38.16	C	
ANISOU	3367	CB	ILE	B	191	4051	4459	5989	255	52	-365	C
ATOM	3368	CG1	ILE	B	191	69.193	-7.003	19.772	1.00	47.02	C	
ANISOU	3358	CG1	ILE	B	191	5185	5594	7085	254	145	-384	C
ATOM	3369	CG2	ILE	B	191	70.375	-3.633	21.325	1.00	46.68	C	
ANISOU	3369	CG2	ILE	B	191	5064	5539	7133	312	15	-389	C
ATOM	3370	CD1	ILE	B	191	70.529	-6.243	19.558	1.00	43.23	C	
ANISOU	3370	CC1	ILE	B	191	4526	5101	6556	240	202	-407	C
ATOM	3371	N	HIS	B	192	68.294	-8.118	25.975	1.00	30.63	N	
ANISOU	3371	N	HIS	B	192	3133	3438	5018	213	-116	-295	N
ATOM	3372	CA	HIS	B	192	68.392	-8	693	25.237	1.00	30.63	C
ANISOU	3372	CA	HIS	B	192	3100	3492	5041	222	-199	-275	C
ATOM	3373	C	HIS	B	192	67.558	-9.509	25.025	1.00	33.40	C	
ANISOU	3373	C	HIS	B	192	3529	3300	5302	233	-254	-2316	C
ATOM	3374	O	HIS	B	192	57.836	-10.532	25.977	1.90	33.51	O	
ANISOU	3374	O	HIS	B	192	3539	3787	5407	275	-312	-220	O
ATOM	3375	CB	HIS	B	192	68.824	-7.592	26.308	1.00	31.59	C	
ANISOU	3375	CB	HIS	B	192	3202	31354	5148	164	-217	-255	C
ATOM	3376	CG	HIS	B	192	69.122	-8.147	27.840	1.00	29.28	C	
ANISOU	3376	CG	HIS	B	192	2389	3775	4362	172	-332	-242	C
ATOM	3377	ND1	HIS	B	192	70.190	-5.901	27.335	1.00	03.48	N	
ANISOU	3377	ND1	HIS	B	192	3750	7.915	5447	220	-342	-250	N
ATOM	3378	CD2	HIS	B	192	65.385	-8.170	28.826	1.00	130.18	C	
ANISOU	3378	CD2	HIS	B	192	3041	3495	4930	144	-300	-203	C
ATOM	3379	OE1	HIS	B	192	70.202	-9.360	29.157	1.00	31.77	O	
ANISOU	3379	OE1	HIS	B	192	3141	3711	5220	222	-429	-210	O
ATOM	3380	NE2	HIS	B	192	69.097	-8.907	29.738	1.00	32.79	N	
ANISOU	3380	NE2	HIS	B	192	3332	3840	5287	174	-4516	-186	N
ATOM	3381	N	ALA	B	193	57.317	-9.205	25.557	1.03	20.441	N	
ANISOU	3381	N	ALA	B	193	3220	3420	4932	193	-236	-216	N
ATOM	3382	CA	ALA	B	193	65.154	-10.046	26.012	1.00	29.44	C	
ANISOU	3382	CA	ALA	B	193	3155	3257	4772	190	-256	-176	C
ATOM	3383	C	ALA	B	193	64.987	-41.265	25.115	1.30	29.91	C	
ANISOU	3383	C	ALA	B	193	3244	3265	4555	2017	-290	-188	C
ATOM	3384	O	ALA	B	193	64.853	-12.425	25.572	1.00	20.56	O	
ANISOU	3384	O	ALA	B	193	3218	3150	4832	201	-349	-164	O
ATOM	3385	CB	ALA	B	193	63.884	-9.225	26.065	1.00	27.81	C	
ANISOU	3385	CB	ALA	B	193	3001	3062	4502	145	-239	-1.57	C
ATOM	3386	N	ALA	B	194	65.054	-11.040	23.820	1.00	23.98	N	
ANISOU	3386	N	ALA	B	194	3127	3147	4738	253	-2796	-227	N

TABLE 3-continued

ATOM	3387	CA	ALA	B	194	65.002	-12.153	22.914	1.00	32.16	C
ANISOU	3387	CA	ALA	B	194	3554	3503	5161	300	-227	-251 C
ATOM	3388	C	ALA	B	194	00.099	-13.158	23.02	1.00	33.05	C
ANISOU	3388	C	ALA	B	194	3740	3703	5456	355	-266	-265 C
ATOM	3389	O	ALA	B	194	65802	-14.410	22.972	1.00	33.95	O
ANISOU	8389	O	ALA	B	194	3774	3648	5431	393	-304	-265 O
ATOM	3390	CB	ALA	B	194	65.005	-11.641	21.480	1.00	32.97	C
ANISOU	3390	CB	ALA	B	194	3651	3023	5244	309	-1413	-294 C
ATOM	3391	N	GLY	B	195	67.321	-12.701	23.332	1.00	31.29	N
ANISOU	3391	N	GLY	B	195	3322	3398	5145	373	-254	-281 N
ATOM	3392	CA	GLY	B	195	65.442	-13.52	23.573	1.00	31.60	C
ANISOU	3392	CA	GLY	B	195	3324	3430	5246	434	-292	-293 C
ATOM	3393	C	GLY	B	195	68.202	-14.457	24.517	1.00	35.22	C
ANISOU	3393	C	GLY	B	195	3794	3854	5733	443	-383	-242 C
ATOM	3394	O	GLY	B	195	56.745	-15.005	24.573	1.00	34.74	O
ANISOU	3394	O	GLY	B	195	3728	3749	5724	503	-425	-242 O
ATOM	3395	N	LEU	B	196	67.591	-13.959	25.831	1.00	33.44	N
AMISOU	3395	N	LEU	B	196	3002	3049	5463	387	-414	-195 N
ATOM	3396	CA	LEU	B	196	57.351	-14.803	27.025	1.00	33.40	C
ANISOU	3396	CA	LEU	B	196	3615	3623	5470	393	-498	-138 C
ATOM	3397	C	LEU	B	196	65.452	-16.051	25.729	1.00	34.14	C
ANISOU	3397	C	LEU	B	196	3778	3029	5555	411	-524	-1113 C
ATOM	3398	O	LEU	B	196	60.747	-17.101	27.246	1.00	32.70	O
ANISOU	3398	O	LEU	B	196	31505	3398	5423	451	-585	-88 O
ATOM	3399	CB	LEU	B	196	66.680	-13.969	28.143	1.00	31.43	C
ANISOU	3399	CB	LEU	B	196	3378	3412	5155	324	-519	-93 C
ATOM	3400	OG	LEU	B	196	67.029	-12.800	28.792	1.00	31.82	O
ANISOU	3400	OG	LEU	B	196	3339	3537	5190	301	-513	-107 O
ATOM	3401	CD1	LEU	B	196	60	577	-11.342	29.542	1.00	29.84 C
ANISOU	3401	CD1	LEU	B	196	3142	3322	4874	231	-510	-81 C
ATOM	3402	CD2	LEU	B	196	68.799	-13.299	29.602	1.00	34.84	C
ANISOU	3402	CD2	LEU	B	196	3674	3939	5626	348	-578	-98 C
ATOM	3403	N	ALA	B	197	55.458	-15.865	25.901	1.00	31.75	N
ANISOU	3403	N	ALA	B	197	3525	a.311	5225	379	-480	-13.5 N
ATOM	3404	CA	ALA	B	197	64.625	-15.995	26.403	1.00	32.72	C
AMISOU	3404	CA	ALA	B	197	3716	3358	5359	389	-497	-133 C
ATOM	3405	C	ALA	B	197	65.430	-17.950	24.512	1.00	31.01	C
ANISOU	4405	C	ALA	B	197	3487	3090	5207	454	-492	-132 C
ATOM	3406	O	ALA	B	197	65.397	-19.196	24.673	1.00	31.92	O
ANISOU	3406	O	ALA	B	197	3634	3128	5300	498	-540	-168 O

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ATOM	3407	CB	ALA	B	197	63.390	-16.441	24.559	1.00	30.29	C
ANISOU	3407	CB	ALA	B	197	3453	3062	4992	338	-450	-147 C
ATOM	3408	N	TYR	B	198	66.112	-17.394	23.540	3.00	39.47	N
ANISOU	3408	N	TYR	B	198	3376	3957	5144	489	-430	-242 N
ATOM	3409	CA	TYR	B	198	67.005	-18.155	22.582	1.00	34.35	C
ANISOU	3409	CA	TYR	B	198	3344	3515	5592	1-365	-415	-293 C
ATOM	3410	C	TYR	B	198	67.960	-39.049	23.458	1.00	86.58	C
ANISOU	3410	C	TYR	B	198	4090	3760	50141	626	-477	-277 C
ATOM	3411	O	TYR	B	198	55.078	-20.252	23.129	1.00	35.44	O
ANISOU	3411	O	TYR	B	198	3975	3541	5949	681	-502	-295 O
ATOM	3412	CB	TYR	B	198	67.761	-17.212	21.747	1.00	33.03	C
ANISOU	3412	CB	TYR	B	198	3725	3519	5621	57	-3445	-354 C
ATOM	3413	CG	TYR	B	198	68.6195	7.930	20.772	1.00	39.44	C
ANISOU	3413	CG	TYR	B	198	4402	4253	6380	557	-308	-418 C
ATOM	3414	CD1	TYR	B	198	68.214	-18.350	19.514	1.00	39.130	C
ANISOU	3414	CD1	TYR	B	198	4495	4219	6436	575	-267	-472 C
ATOM	3415	CD2	TYR	B	198	70.042	-130385	21.104	1.00	39.29	C
ANISOU	3415	CD2	TYR	B	198	4307	4198	6425	717	-322	-429 C
ATOM	3416	CE1	TYR	B	198	69.047	-15.964	18.589	1.00	43.17	C
ANISOU	3416	CE1	TYR	B	198	4896	4630	6875	750	-235	-538 C
ATOM	3417	CE2	TYR	B	198	70.873	-13.652	20.138	1.00	44.26	C
ANISOU	3417	CE2	TYR	B	198	4906	4810	7102	797	-292	-493 C
ATOM	3418	CZ	TYR	B	198	70.331	-19.245	18.940	1.00	40.95	C
ANISOU	3418	CZ	TYR	B	198	4541	4358	0659	811	-247	-547 C
ATOM	3419	OH	TYR	B	198	71.114	-19.821	1.5014	1.00	46.84	O
ANISOU	3419	OH	TYR	B	198	5514	5347	7597	588	-211	-615 O
ATOM	3420	N	LYS	B	199	58.588	-18.	517	24.510	1.00	35.36 N
ANISOU	3420	N	LYS	B	199	38813	3659	5892	611	-507	-2139 N
ATOM	3421	CA	LYS	B	199	69.623	-19.275	25.227	1 00	39.91	C
ANISOU	3421	CA	LYS	B	199	4414	4218	0533	684	-557	-220 C
ATOM	3422	C	LYS	B	199	69.003	-20.371	26.050	1.00	43.57	C
ANISOU	3422	C	LYS	B	199	4942	4505	7005	588	-643	-156 C
ATOM	3423	O	LYS	B	199	69.085	-21.260	25.504	1.50	44.13	O
ANISOU	3423	O	LYS	B	199	4998	46.33	7138	753	-696	-138 O
ATOM	3424	CB	LYS	B	199	19.410	-13.387	25.130	1.00	41.15	C
ANISOU	3424	CB	LYS	B	199	4495	4458	5681	606	-585	-197 C
ATOM	3425	CB	LYS	B	199	71.353	-17.444	25.435	1.00	43.13	C
ANISOU	3425	CB	LYS	B	199	4064	4760	6943	673	-516	-258 C
ATOM	3426	CD	LYS	B	199	71.595	-16.348	26.350	1.00	50.45	C
ANISOU	3426	CD	LYS	B	199	5525	5792	7851	630	-528	-249 C

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ATOM	3427	CE	LYS	B	199	72.657	-15.315	25.509	1.00	57.34	C
ANISOU	3427	CE	LYS	B	199	8301	6703	8705	618	-446	-390 C
ATOM	3428	NZ	LYS	B	199	73.926	-14.846	25.133	140	44.58	N
ANISOU	3428	NZ	LYS	B	199	7133	7715	9664	579	-466	-308 N
ATOM	3429	N	ARG	B	200	67.691	-201.282	25.246	1.00	38.18	N
ANISOU	3429	N	ARG	B	200	4331	2501	6273	6119	-546	-121 N
ATOM	3430	CA	ARG	B	200	66.972	-21.325	25.958	1.00	38.04	C
ANISOU	3430	CA	ARG	B	200	4332	13807	5255	611	-710	-57 C
ATOM	3431	C	ARG	B	200	65.303	-22.326	26.084	1.00	37.24	C
ANISOU	3431	C	ARG	B	200	4346	3510	6193	523	-702	-65 C
ATOM	3432	O	ARG	B	200	65.454	-23.568	25.577	1.00	40.75	O
ANISOU	3432	O	ARG	B	200	4858	3992	6638	594	-744	-33 O
ATOM	3433	CB	ARG	B	200	05.931	-20.734	27.900	1.00	37.92	C
ANISOU	3433	CB	ARG	B	200	4400	3629	6179	527	-726	7 C
ATOM	3434	CG	ARG	B	200	66.520.	-20.110	29.133	1 00	33.72	C
ANISOU	3434	CG	ARG	B	200	5085	4637	6891	519	-763	513 C
ATOM	3435	CD	ARG	B	200	65.432	-19.282	29.843	1.00	41.95	C
ANISOU	3435	CD	ARG	B	200	4891	4454	6585	431	-757	96 C
ATOM	3436	NE	ARG	B	200	64.304	-20.034	30.463	1.00	43.84	N
ANISOU	3436	NE	ARG	B	200	4824	4263	6-1426	396	-797	166 N
ATOM	3437	CZ	ARG	B	200	64.414	-23.710	31.148	1.00	45.47	C
ANISOU	3437	CZ	ARG	B	200	5425	4839	7012	404	-8.64	244 C
ATOM	3438	NH1	ARG	B	200	65.522	-20.641	132.a68	1.00	44.36	N
ANISOU	3438	NH1	ARG	B	200	5234	4800	6950	450	-306	259 N
ATOM	3439	NH2	ARG	B	200	63.3934	-21.422	32.167	1.03	41.48	N
ANISOU	3439	NH2	ARG	B	200	4937	4234	6493	366	-92	310 N
ATOM	3440	N	ASP	B	201	66.615	-22.355	24.787	1.00	35.72	N
ANISOU	3440	N	ASP	B	201	44 3410	6019	6t.0	-648	-168	N
ATOM	3441	CA	ASP	B	201	66.078	-23.351	23.846	1.00	37.86	C
ANISOU	3441	CA	ASP	B	201	4477	3533	631	577	-641	-211 C
ATOM	3442	C	ASP	B	201	64.532	-23.107	23.606	1.00	33.51	C
ANISOU	3442	C	ASP	B	201	4750	3796	6466	590	-526	-201 C
ATOM	3443	O	ASP	B	201	63.829	-24.043	23.404	1.00	37.54	O
ANISOU	3443	O	ASP	B	201	4564	3462	6206	577	-650	-200 O
ATOM	3444	CB	ASP	B	201	66.344	-24813	24.316	1.00	40.69	C
ANISOU	3444	CB	ASP	B	201	4870	3835	6754	733	-708	-181 C
ATOM	3445	CG	ASP	B	201	65003	-25.916	23.132	1.33	45.82	C
ANISOU	3445	CG	ASP	B	201	5706	4507	7575	765	-697	-240 C
ATOM	3446	OD1	ASP	B	201	65.992	-25.637	21.343	1.00	44.27	O
ANISOU	3446	OD1	ASP	B	201	5379	4206	7234	774	-637	-334 O

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ATOM	3447	OD2	ASP	B	201	65.701	-27.038	23.564	1.30	47.52	O	
ANISOU	3447	OD2	ASP	B	201	5853	4487	77153	773	-750	-215	O
ATOM	3448	N	ILE	B	202	64.162	-21.840	23.565	1.00	36.68	N	
ANISOU	3448	N	ILE	B	202	4397	3555	6062	531	-534	-1313	N
ATOM	3449	CA	ILE	B	202	62.772	-21.461	23.367	1.30	35.51	C	
ANISOU	3449	CA	ILE	B	202	4271	3394	5825	455	-558	-18	C
ATOM	3450	C	ILE	B	202	62.927	-20.912	21.940	1.00	37.17	C	
ANISOU	3450	C	ILE	B	202	4471	3642	6010	468	-428	-269	C
ATOM	3451	O	ILE	B	202	63.755	-20.753	21.532	1.00	37.24	O	
ANISOU	3451	O	ILE	B	202	4423	3739	6316	5.040	-455	-302	O
ATOM	3452	CB	ILE	B	202	52.243	-23.434	24.390	1.00	35.32	C	
ANISOU	3452	CB	ILE	B	202	4311	3520	5818	389	-572	-123	C
ATOM	3453	CG1	ILE	B	202	62.396	-21022	25.833	1.00	30.15	C	
ANISOU	3453	CG1	ILE	B	202	3366	3143	5490	2851	-642	-42	C
ATOM	3454	CG2	ILE	B	202	60.807	-13.933	24.326	1.00	34.84	C	
ANISOU	3454	CG2	ILE	B	202	4216	3404	5617	315	-548	-121	C
ATOM	3455	CD1	ILE	B	202	51.525	-2230	26.043	1.00	40.86	C	
ANISOU	3455	CD1	ILE	B	202	5008	4019	6437	373	-683	-6	C
ATOM	3456	N	PRO	B	203	63.836	-21.311	21.097	1.00	26.92	N	
ANISOU	3456	N	PRO	B	203	4490	3577	5960	446	-434	-304	N
ATOM	3457	CA	PRO	B	203	61.961	-20.762	19750	1.30	33.26	C	
ANISOU	3457	CA	PRO	B	203	4017	3153	5461	462	-422	-374	C
ATOM	3458	C	PRO	B	203	61.770	-19.211	19.752	1.00	34.15	C	
ANISOU	3458	C	PRO	B	203	4099	3374	5533	420	-372	-361	C
ATOM	3459	O	PRO	B	203	60.319	-10.713	20.475	1.30	32.93	O	
ANISOU	3459	O	PRO	B	203	3964	3250	5316	263	-338	-307	O
ATOM	3460	CB	PRO	B	203	60.851	-21.431	9.315	1.00	37.47	C	
ANISOU	3460	CB	PRO	B	203	4665	3695	6029	434	-431	-410	C
ATOM	3461	CG	PRO	B	203	63.120	-22.309	23.326	1.00	301.74	C	
ANISOU	3461	CG	PRO	B	203	4343	3863	5297	7.93	-499	-343	C
ATOM	3462	CD	PRO	B	203	60.775	-22.250	21.313	1.013	43.28	C	
ANISOU	3462	CD	PRO	B	203	4932	3928	7396	434	-534	-279	C
ATOM	3463	N	VAL	B	204	52.566	-13.482	18.050	1.00	34.10	N	
ANISOU	3463	N	VAL	B	204	4051	3422	5483	453	-313	-436	N
ATOM	3464	CA	VAL	B	204	62.480	-17.052	18.852	1.00	32.36	C	
ANISOU	3464	CA	VAL	B	204	3883	3362	5282	419	-259	-393	C
ATOM	3465	C	VAL	B	204	52.402	-16.796	17.369	1.33	32.79	C	
ANISOU	3465	C	VAL	B	204	3869	3368	5222	439	-197	-456	C
ATOM	3466	O	VAL	B	204	62.337	-17.165	15.605	1.00	03.31	O	
ANISOU	3466	O	VAL	B	204	3912	3428	5316	498	-167	-511	O

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ATOM	3467	CB	VAL	B	204	63.738	-15.38	19.446	1.00	31.23	C
ANISOU	3467	CB	VAL	B	204	3590	3155	5091	438	-243	-380 C
ATOM	3468	CG1	VAL	B	204	63.567	-14.880	19.375	1.00	33.15	C
ANISOU	3468	CG1	VAL	B	204	3434	3122	4899	395	-190	-365 C
ATOM	3469	CG2	VAL	B	204	53.940	-16.726	20.919	1.00	27.58	C
ANISOU	3469	CG2	VAL	B	204	3115	2599	4564	429	-311	-321 C
ATOM	3470	N	VAL	B	205	61.327	-16.140	16.986	1.00	39.29	N
ANISOU	3470	N	VAL	B	205	3572	3075	4826	394	-176	-446 N
ATOM	3471	CA	VAL	B	205	61.302	-15.788	15.595	1.00	30.77	C
ANISOU	3471	CA	VAL	B	205	3651	3185	4840	408	-1.20	-496 C
ATOM	3472	C	VAL	B	205	51.325	-14.264	15.500	1.00	131.87	C
ANISOU	3472	C	VAL	B	205	3769	3400	4941	389	-56	-471 C
ATOM	3473	O	VAL	B	205	60.760	-13.457	16.256	3.00	34.93	O
ANISOU	3473	O	VAL	B	205	4156	3810	5304	342	-66	-418 O
ATOM	3474	CB	VAL	B	205	59.677	-15.134	1.5.124	1.00	34.10	C
ANISOU	3474	CB	VAL	B	205	4141	3595	5220	375	-142	-503 C
ATOM	3475	CG1	VAL	B	205	59.511	-15.710	12.66.5	1.00	34.14	C
ANISOU	3475	CG1	VAL	B	205	4159	3650	5161	3915	-86	-554 C
ATOM	3476	CG2	VAL	B	205	69.349	-17.527	15.350	1.00	32.90	C
ANISOU	3476	CG2	VAL	B	205	4024	3358	5118	379	-209	-521 C
ATOM	3477	N	SER	B	206	52.209	-13.854	14.607	1.00	34.14	N
ANISOU	3477	N	SER	B	206	4026	2721	5220	426	6	-508 N
ATOM	3478	CA	SER	B	206	62.429	-12.412	14.425	1.09	33.93	C
ANISOU	3478	CA	SER	B	206	3976	3759	5156	404	70	-4614 C
ATOM	3479	C	SER	B	206	61.664	1.879	12.243	1.00	39.67	C
ANISOU	3479	C	SER	B	206	3601	3387	4664	401	117	-496 C
ATOM	3480	O	SER	B	206	61.737	-12.485	12.151	1.00	35.23	O
ANISOU	3480	O	SER	B	206	4497	2956	5221	443	135	-550 O
ATOM	3481	CB	SER	B	206	53.933	-12.100	14.270	1.00	39.12	C
ANISOU	3481	CB	SER	B	206	4569	4436	5857	440	117	-506 C
ATOM	3482	OG	SER	B	206	54.453	-12.340	15.603	1.00	43.35	O
ANISOU	3482	OG	SER	B	206	5068	4949	6455	429	67	-475 O
ATOM	3483	N	LEU	B	207	61.037	-10.703	13.394	1.00	30.04	N
ANISOU	3483	N	LEU	B	207	2531	13344	4537	352	140	-451 N
ATOM	3484	CA	LEU	B	207	50.406	-10.034	121199	1.00	30.81	C
ANISOU	3484	CA	LEU	B	207	3653	3489	4555	366	392	-455 C
ATOM	3485	C	LEU	B	207	61.141	-8.742	11.905	1.00	31.29	C
ANISOU	3485	C	LEU	B	207	3693	3591	4632	362	269	-433 C
ATOM	3486	O	LEU	B	207	60.993	-7.850	13.717	1.00	32.74	O
ANISOU	3486	O	LEU	B	207	3868	3778	4793	325	270	-387 O

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ATOM	3487	CB	LEU	B	207	58.918	-9.594	12.497	1.00	23.26	C
ANISOU	3487	CB	LEU	B	207	4027	3520	4530	227	157	-417 C
ATOM	3488	CG	LEU	B	207	57.927	-10.344	12.811	1.00	33.86	C
ANISOU	3488	CG	LEU	B	207	4121	3546	4599	313	80	-429 C
ATOM	3489	CD1	LEU	B	207	56.458	-10.423	12.854	1.00	35.46	C
ANISOU	3489	CD1	LEU	B	207	4355	4074	5044	280	58	-395 C
ATOM	3490	CD2	LEU	B	207	58.032	-12.106	11.968	1.00	37.58	C
ANISOU	3490	CC2	LEU	B	207	4612	4291	5377	346	61	-404 C
ATOM	3491	N	THR	B	208	61.900	-8.618	10.810	1.00	36.47	N
ANISOU	3491	N	THR	B	208	338	4279	5243	397	236	-455 N
ATOM	3492	CA	THR	B	208	52.918	-7.567	10	630	1.00	33.75 C
ANISOU	3492	CA	THR	B	208	3952	3967	4906	393	413	-447 C
ATOM	3493	C	THR	B	208	62.758	-7.046	9.239	1.00	32.20	C
ANISOU	3493	C	THR	B	208	2784	13819	4633	413	434	-452 C
ATOM	3494	O	THR	B	208	61.922	-7.544	3.436	1.00	38.97	O
ANISOU	3494	O	THR	B	208	3676	3677	441	5	435	469 -474 O
ATOM	3495	CB	THR	B	208	64.370	-8.160	10.759	1.00	37.79	C
ANISOU	3495	CB	THR	B	208	4389	4469	54610	422	427	-467 C
ATOM	3496	OG1	THR	B	208	64.605	-9.097	9.662	1.00	06.26	O
ANISOU	3496	OG1	THR	B	208	4473	4541	558.2	477	444	-543 O
ATOM	3497	OG2	THR	B	208	64.456	-9.012	1.931	1.00	36.13	O
ANISOU	3497	OG2	THR	B	208	4171	4209	5348	47	344	-490 O
ATOM	3498	N	ASN	B	209	83.552	-6.045	8	931	1.30	31.22 N
ANISOU	3498	N	ASN	B	209	3630	3723	4503	405	562	-429 N
ATOM	3499	CA	ASN	B	209	83.647	-5.6019	7.554	1.00	32.39	C
ANISOU	3499	CA	ASN	B	209	3303	3966	4649	409	640	-432 C
ATOM	3500	C	ASN	B	209	613.063	-5.720	7.093	1.00	35.06	C
ANISOU	3500	C	ASN	B	209	4330	4284	4953	453	706	-464 C
ATOM	3501	O	ASN	B	209	65.508	-4.904	6.191	1.00	04.98	O
ANISOU	3501	O	ASN	B	209	4067	4316	4906	458	792	-449 O
ATOM	3502	CB	ASN	B	209	63.197	-4.126	7.436	1.00	31.33	C
ANISOU	3502	CB	ASN	B	209	3753	3847	4472	396	589	-363 C
ATOM	3503	CG	ASN	B	209	61.716	-8.938	7.774	1.00	34.03	C
ANISOU	3503	CG	ASN	B	209	4087	4136	4706	381	631	-330 C
ATOM	3504	OD1	ASN	B	209	60.910	-3.576	6.923	1.00	136.65	O
ANISOU	3504	OD1	ASN	B	209	4467	4500	4960	396	643	-311 O
ATOM	3505	ND2	ASN	B	209	61.379	-4.120	9.004	1.00	30.66	N
ANISOU	3505	ND2	ASN	B	209	3658	3667	4331	352	565	-321 N
ATOM	3506	N	ILE	B	210	65.833	-5.611	7.702	1.00	34.73	N
ANISOU	3506	N	ILE	B	210	13995	4221	4998	459	569	-505 N

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ATOM	3507	CA	ILE	B	210	67.227	-9.753	7.272	1.00	35.05	C	
ANISOU	3507	CA	ILE	B	210	4392	4410	5196	496	733	-540 C	
ATOM	3508	C	ILE	B	210	67.632	-8.221	7.514	1.00	36.23	C	
ANISOU	3508	C	ILE	B	210	4345	4655	5526	545	676	-685 C	
ATOM	3509	O	ILE	B	210	67.224	-5.304	4.521	1.00	3459	O	
ANISOU	3509	O	ILE	B	210	3691	4143	5100	535	590	-602 O	
ATOM	3510	CB	ILE	B	210	68.151	-5.754	8.040	1.00	36.14	C	
ANISOU	3510	CB	ILE	B	210	4039	4422	5272	449	768	-501 C	
ATOM	3511	CG1	ILE	B	210	69.568	-5.758	7.459	1.00	4149	C	
ANISOU	3511	CG1	ILE	B	210	4691	5193	6831	473	846	-532 C	
ATOM	3512	CG2	ILE	B	210	68.184	-6.087	9.521	1.00	38.13	C	
ANISOU	3512	CG2	ILE	B	210	4007	4372	5347	42.6	682	-496 C	
ATOM	3513	CD1	ILE	B	210	69.631	-5.909	6.145	1.00	42.07	C	
ANISOU	3513	CD1	ILE	B	210	47135	5271	5977	476	951	-1315 C	
ATOM	3514	N	ASN	B	211	68.438	-3.826	6.641	1.30	37.22	N	
ANISOU	3514	N	ASN	B	211	4191	4555	5396	599	723	-662 N	
ATOM	3515	CA	ASN	B	211	63.767	-13.262	6.837	1.00	42.30	C	
ANISOU	3515	CA	ASN	B	211	4819	5160	6092	653	667	-727 C	
ATOM	3516	C	ASN	B	211	89.763	-10.527	7.962	1.00	44.21	C	
ANISOU	3516	C	ASN	B	211	4934	5374	8440	654	633	-726 C	
ATOM	3517	O	ASN	B	211	69.606	-11.496	6.679	1.00	42.79	O	
ANISOU	3517	O	ASN	B	211	4810	5138	6310	673	552	-743 O	
ATOM	3518	CB	ASN	B	211	69.249	-18.919	5.519	1.00	49.36	C	
ANISOU	3518	CB	ASN	B	211	5776	6156	7012	723	724	-789 C	
ATOM	3519	CG	ASN	B	211	63.100	-11.074	4.537	1.00	56.09	C	
ANISOU	3519	CG	ASN	B	211	66.54	6961	7697	729	723	-817 C	
ATOM	3520	OD1	ASN	B	211	87.911	-12.002	4.63	2133	1.00	60.74 O	
ANISOU	3520	OD1	ASN	B	211	7292	7504	3284	742	649	-849 O	
ATOM	3521	ND2	ASN	B	211	67.981	-10.156	3.539	1.00	54.42	N	
ANISOU	3521	ND2	ASN	B	211	6462	91817	7400	720	803	-793 N	
ATOM	3522	N	HIS	B	212	791.7819	-9.540	8.425	1.00	40.35	N	
ANISOU	3522	N	HIS	B	212	4426	4929	5988	631	594	-702 N	
ATOM	3523	CA	HIS	B	212	71.821	-9.8610	0.125	1.00	39.54	C	
ANISOU	3523	CA	HIS	B	212	4234	430	5980	536	564	-706 C	
ATOM	3524	C	HIS	B	212	71.922	-3.729	10.039	1.00	21771	C	
ANISOU	3524	C	HIS	B	212	3972	458.3	5773	564	1562	-647 C	
ATOM	3525	O	HIS	B	212	72.379	-7-558	9.599	1.00	37.72	O	
ANISOU	3525	O	HIS	B	212	3954	4524	5743	522	739	-9116 O	
ATOM	3526	CB	HIS	B	212	73.162	-9.9781	8.447	1.00	41.13	C	
ANISOU	3526	CB	HIS	B	212	4354	5366.	32-111	1	680	741	-760 C

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TABLE 3-continued

ATOM	3527	CG	HIS	B	212	73.223	-10931	7.392	1.00	43	90	C
ANISOU	3527	CG	HIS	B	212	4732	1423	6524	754	770	-817	C
ATOM	3528	ND1	HIS	B	212	73.223	-12.2518	7.479	1.00	50.73	N	
ANISOU	3528	ND1	HIS	B	212	5511	5246	7437	817	704	-670	N
ATOM	3529	CD2	HIS	B	212	73.108	-10.714	5.9132	1.00	45.88	C	
ANISOU	3529	CD2	HIS	B	212	5011	5727	6694	775	855	-841	C
ATOM	3530	CE1	HIS	B	212	73.222	-12.881	6.278	1.00	46.31	C	
ANISOU	3530	CE1	HIS	B	212	5074	5700	6823	876	751	-930	C
ATOM	3531	NE2	HIS	B	212	78.142	-11.946	5.338	1.05	49.86	N	
ANISOU	3531	NE2	HIS	B	212	55136	5216	7193	851	841	-916	N
ATOM	3532	N	SER	B	213	71.910	-9.0158	11.310	1.09	35.75	N	
ANISOU	3532	N	SER	B	213	3700	4294	5582	551	576	-531	N
ATOM	3533	CA	SER	B	213	71.899	-8.052	12.387	1.00	35.09	C	
ANISOU	3533	CA	SER	B	213	3600	42021	5522	481	556	-580	C
ATOM	3534	C	SER	B	213	72.253	-8.077	13.647	1.00	32.30	C	
ANISOU	3534	C	SER	B	213	3283	3897	5316	498	4611	-565	C
ATOM	3535	O	SER	B	213	72.303	-10.100	10.577	1.90	3480	O	
ANISOU	3535	O	SER	B	213	3580	4110	5580	559	417	-518	O
ATOM	3536	CB	SER	B	213	70.441	-7534	12.534	1.00	35.54	C	
ANISOU	3536	CB	SER	B	213	8750	4241	5513	435	531	-536	C
ATOM	3537	OG	SER	B	213	69.431	-8.637	13.025	1.04	35.38	O	
ANISOU	3537	OG	SER	B	213	3779	4171	5493	451	441	-543	O
ATOM	3538	N	PRO	B	214	72.549	-8.228	14.791	1.00	33.80	N	
ANISOU	3538	N	PRO	B	214	33.58	4016	5450	448	427	-553	N
ATOM	3539	CA	PRO	B	214	72.781	-8.926	16.081	1.00	37.95	C	
ANISOU	3539	CA	PRO	B	214	38137	4516	6050	451	332	-547	C
ATOM	3540	C	PRO	B	214	71.657	-3.848	16.432	1.00	37.64	C	
ANISOU	3540	C	PRO	B	214	3697	4418	5986	478	253	-533	C
ATOM	3541	O	PRO	B	214	71.900	-10.332	17.066	1.00	35.65	O	
ANISOU	3541	O	PRO	B	214	3759	4261	.5905	519	182	-539	O
ATOM	3542	CB	PRO	B	214	72.862	-7.782	174342	1.00	86.22	C	
ANISOU	3542	CB	PRO	B	214	8608	4314	5640	386	320	-511	C
ATOM	3543	CG	PRO	B	214	73.661	-6.783	15.233	1.00	35.43	C	
ANISOU	3543	CG	PRO	B	214	3582	4855	5870	861	421	-523	C
ATOM	3544	CD	PRO	B	214	72.855	-6.730	14.952	1.00	34.84	C	
ANISOU	3544	CD	PRO	B	214	3714	4437	5855	378	481	-525	C
ATOM	3545	N	LEU	B	215	70.430	-9.449	16.150	1.09	33.89	N	
ANISOU	3545	N	LEU	B	215	3504	3929	5424	440	262	-.509	N
ATOM	3546	CA	LEU	B	215	69.299	-10.299	16.524	1.00	05.481	C	
ANISOU	3646	CA	LEU	B	215	3779	4078	5524	450	139	-404	C

TABLE 3-continued

ATOM	3547	C	LEU	B	215	392317	-11.517	15.725	1.00	86.16	C
ANISOU	3547	C	LEU	B	215	3892	4181	5717	518	179	-539
ATOM	3548	O	LEU	B	215	68.737	-12.558	16.205	1.00	34.29	O
ANISOU	3548	O	LEU	B	215	3590	3839	5496	539	106	-537
ATOM	3549	CB	LEU	B	215	67.984	-9.430	16.319	1.00	31.19	C
ANISOU	3549	CB	LEU	B	215	3309	3537	.5004	393	207	-461
ATOM	3550	CG	LEU	B	215	66.6167	-10.220	10.500	1.00	33.41	C
ANISOU	3550	CG	LEU	B	215	3665	8774	5254	395	144	-445
ATOM	3551	CD1	LEU	B	215	65.534	-10.539	18.015	1.00	33.42	C
ANISOU	3551	CD1	LEU	B	215	3552	3746	5290	377	50	5011
ATOM	3552	CD2	LEU	B	215	65.537	-9.253	13.136	1.30	32.161	C
ANISOU	3552	CD2	LEU	B	215	3553.	3635	5022	352	179	-418
ATOM	3553	N	LEU	B	215	69.759	-11.611	14.507	1.00	59.04	N
ANISOU	3553	N	SER	B	216	4239	4525	15070	5.54	252	-5.84
ATOM	3554	CA	SER	B	216	69.730	-12.870	13.789	3.30	41.54	C
ANISOU	3554	CA	SER	B	216	4.584	4505	6592	519	243	-535
ATOM	3555	C	SER	B	216	70.510	-14.040	14.439	1.00	43.26	C
ANISOU	3555	C	SER	B	216	4762	4931	51592	577	178	-555
ATOM	3556	O	SER	B	216	70.053	-35.185	14.344	1400	43.51	O
ANISOU	3556	O	SER	B	216	4341	4254	6737	714	131	-677
ATOM	3557	CB	SER	B	216	59.893	-12.729	12.279	1.00	46.49	C
ANISOU	3557	CB	SER	B	216	5220	5473	5971	549	327	-682
ATOM	3558	CG	SER	B	216	11.077	-12.142	13.898	1.00	44.461	C
ANISOU	3558	OG	SER	B	216	6	4855	5271	57155	6512	4010
ATOM	3559	N	SER	B	217	71.550	-13	7515	15.204	1.00	38.07
ANISOU	3559	N	SER	B	217	4023	4349	5092	673	159	-541
ATOM	3560	CA	SER	B	217	72.286	-14.804	15.912	1.00	411.96	C
ANISOU	3560	CA	SER	B	217	4475	4805	6663	737	104	-650
ATOM	3561	C	SER	B	217	71.513	-15.3135	17.087	1.00	41.25	C
ANISOU	3561	C	SER	B	217	4424	46519	9552	714	9	-503
ATOM	3562	O	SER	B	217	71.970	-16.355	17.332	1.510	35.97	O
ANISOU	3562	O	SER	B	217	3863	4372	6092	755	-57	-591
ATOM	3563	CB	SER	B	217	73.556	-14.165	36.544	1.00	43.28	C
ANISOU	3563	CB	SER	B	217	4534	5030	5582	733	115	-641
ATOM	3564	OG	SER	B	217	74.502	-14.071	15.552	1.00	55.36	O
ANISOU	3564	OG	SER	B	217	6004	6605	6427	775	194	-692
ATOM	3565	N	LEU	B	218	70.375	-14.558	17.394	3.330	375.25	N
ANISOU	3565	N	LEU	B	218	3951	4153	6015	644	3	-553
ATOM	3566	CA	LEU	B	218	59.503	-14.995	18.590	1.00	36.55	C
ANISOU	3566	CA	LEU	B	218	3940	4020	5927	510	-56	-502

TABLE 3-continued

ATOM	3567	C	LEU	B	218	63.151	-15.365	13.136	1.00	37.55	C
ANISOU	3567	C	LEU	B	218	4163	4104	6002	584	-99	-495 C
ATOM	3568	O	LEU	B	218	57.229	-16.254	18.951	1.00	34.56	O
ANISOU	3568	O	LEU	B	218	3326	3103	5528	535	-144	-447 O
ATOM	3569	CB	LEU	B	218	69.554	-13.759	19.528	1.00	35.00	C
ANISOU	3569	CB	LEU	B	218	3717	3575	5709	541	-551	-455 C
ATOM	3570	CG	LEU	B	218	70.572	-13.347	20.200	1.00	38.56	C
ANISOU	3570	CG	LEU	B	218	4071	4359	5211	550	-94	-454 C
ATOM	3571	CD1	LEU	B	218	70.712	-12.024	23.956	1.00	36.92	C
ANISOU	3571	CD1	LEU	B	218	3843	4209	5975	476	-87	-421 C
ATOM	3572	CD2	LEU	B	218	71.411	4.425	21.329	1.00	40.31	C
ANISOU	3572	CD2	LEU	B	218	4257	4556	5402	600	-178	-439 C
ATOM	3573	N	SER	B	219	57.992	-15.793	16.940	1.00	35.30	N
ANISOU	3573	N	SER	B	219	3907	3807	5700	618	-57	-553 N
ATOM	3574	CA	SER	B	219	56.686	-16.136	15.425	1.30	34.49	C
ANISOU	3574	CA	SER	B	219	3837	3572	5547	594	-57	-554 C
ATOM	3575	C	SER	B	219	56.787	-17.565	15.9.58	1.00	33.05	C
ANISOU	3575	C	SER	B	219	4355	4053	603.9	555	-95	-1403 C
ATOM	3576	O	SER	B	219	67.779	-13.038	15.383	1.00	35.95	O
ANISOU	3576	O	SER	B	219	45163	13785	5813	721	-70	-655 O
ATOM	3577	CB	SER	B	219	66.393	-15.510	15.223	1.00	37.71	C
ANISOU	3577	CB	SER	B	219	4307	4133	5587	581	11	-583 C
ATOM	3578	OG	SER	B	219	66.451	-13.985	15.534	1.00	37.01	O
ANISOU	3578	OG	SER	B	219	4190	4101	5772	531	42	-541 O
ATOM	3579	N	THR	B	220	55.727	-18.269	15.194	1.00	34.80	N
ANISOU	3579	N	THR	B	220	4020	3582	5520	530	-150	-583 N
ATOM	3580	CA	THR	B	220	65.650	-19.509	15.660	1.00	42.71	C
ANISOU	3580	CA	THR	B	220	5062	4508	6558	579	-175	-640 C
ATOM	3581	C	THR	B	220	65.025	-19.583	14.233	1.00	38.77	C
ANISOU	3581	C	THR	B	220	4606	4028	5096	679	-128	-704 C
ATOM	3582	O	THR	B	220	65.253	-20.448	13.415	1.00	40.50	O
ANISOU	3582	O	THR	B	220	4845	4205	6334	731	-121	-773 O
ATOM	3583	CB	THR	B	220	64.916	-20.374	16.743	1.00	35.89	C
ANISOU	3583	CB	THR	B	220	4242	3570	5324	647	-260	-585 C
ATOM	3584	OG1	THR	B	220	65.338	-21.728	16.706	1.00	50.97	O
ANISOU	3584	OG1	THR	B	220	62.98	5519	7933	708	-299	-617 O
ATOM	3585	CG2	THR	B	220	63.612	-20.203	16.575	1.00	25.11	C
ANISOU	3585	CG2	THR	B	220	4199	3457	5673	587	-270	-572 C
ATOM	3586	N	GLU	B	221	64.175	-18.590	13.958	1.00	36.91	N
ANISOU	3586	N	GLU	B	221	4390	31-00	5784	621	-99	-631 N

TABLE 3-continued

ATOM	3587	CA	GLU	B	221	63.530	-16.331D	12.543	1.00	35.05	C
ANISOU	3587	CA	GLU	B	221	4193	3650	5475	618	-55	-731 C
ATOM	3588	C	GLU	B	221	63.477	-16.907	12.344	1.00	34.13	C
ANISOU	3588	C	GLU	B	221	4052	3624	5291	586	9	-701 C
ATOM	3589	O	GLU	B	221	63.543	-16.106	13.277	1.00	34.62	O
ANISOU	3589	O	GLU	B	221	4086	3709	5360	548	3	-638 O
ATOM	3590	CB	GLU	B	221	62.09	-18.900	12.637	1.00	36.89	C
ANISOU	3590	CB	GLU	B	221	4493	3844	5678	570	-107	-726 C
ATOM	3591	CG	GLU	B	221	61.953	-20.352	13.371	1.00	39.35	C
ANISOU	3591	CG	GLU	B	221	4840	4053	0.062	5185	-177	-742 C
ATOM	3592	CD	GLU	B	221	62.338	-2.302	12.026	1.00	47.96	C
ANISOU	3592	CD	GLU	B	221	5950	5100	7173	649	-165	-836 C
ATOM	3593	OE1	GLU	B	221	63.009	-21.013	31.322	1.00	45.18	O
ANISOU	3593	OE1	GLU	B	221	5575	4801	5790	695	-100	-892 O
ATOM	3594	OE2	GLU	B	221	62.068	-22.544	12.245	1.00	49.61	O
ANISOU	3594	OE2	GLU	B	221	6200	5215	7434	650	-222	-855 O
ATOM	3595	N	MET	B	222	63.361	-16.508	11.075	1.00	34.46	N
ANISOU	3595	N	MET	B	222	4108	3719	5266	601	69	-743 N
ATOM	3596	CA	MET	B	222	63.348	-15.052	10.725	1.00	39.94	C
ANISOU	3596	CA	MET	B	222	4784	4495	5397	573	136	-708 C
ATOM	3597	C	MET	B	222	62.376	-14.853	9.624	1.00	40.79	C
ANISOU	3597	C	MET	B	222	4943	4640	5017	565	158	-731 C
ATOM	3598	O	MET	B	222	62.345	-15.64	8.712	1.00	3.9.77	O
ANISOU	3598	O	MET	B	222	4341	41-27	2771	604	159	-831 O
ATOM	3599	CB	MET	B	222	64.686	-14.515	10.182	1.30	43.83	C
ANISOU	3599	CB	MET	B	222	0218	5036	0400	614	214	-731 C
ATOM	3600	CG	MET	B	222	65.833	-14.471	11.185	1.00	55.66	C
ANISOU	3600	CG	MET	B	222	6647	5523	7980	623	207	-708 C
ATOM	3601	SD	MET	B	222	56.516	-12.772	11.295	1.00	71.54	S
ANISOU	3601	SD	MET	B	222	8599	8611	3973	587	235	-659 S
ATOM	3602	CE	MET	B	222	66.460	-12.247	9.589	1.00	51.04	C
ANISOU	3602	CE	MET	B	222	6025	6080	7289	608	381	-555 C
ATOM	3603	N	LEU	B	223	61.534	-13.777	9.708	1.00	36.62	N
ANISOU	3603	N	LEU	B	223	442	4154	5330	518	171	-677 N
ATOM	3604	CA	LEU	B	223	60.899	-12.31313	8.523	1.00	33.25	C
ANISOU	3604	CA	LEU	B	223	4673	4419	5444	520	210	-693 C
ATOM	3605	C	LEU	B	223	61.400	-11.924	8.291	1.00	39.24	C
ANISOU	3605	C	LEU	B	223	4769	4605	5536	514	287	-651 C
ATOM	3606	O	LEU	B	223	61.492	-11.132	9.213	1.00	32.42	O
ANISOU	3606	O	LEU	B	223	3881	3739	4594	478	287	-592 O

TABLE 3-continued

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ATOM	3607	CB	LEU	B	223	55.383	13.291	8.681	1.30	43.51	C	
ANISOU	3607	CB	LEU	B	223	5384	5032	5354	475	157	-657	C
ATOM	3608	CG	LEU	B	223	58.744	-14.712	8.591	1.00	41.95	C	
ANISOU	3608	CG	LEU	B	223	5225	4833	5892	473	86	-721	C
ATOM	3609	CD1	LEU	B	223	57.313	-141570	8.9151	1.00	41.33	C	
ANISOU	3609	CD1	LEU	B	223	5175	4752.	5773	420	30	-584	C
ATOM	3610	CD2	LEU	B	223	58.857	-15.408	7.251	1.00	45.88	C	
ANISOU	3610	CD2	LEU	B	223	5748	5338	5345	523	104	-807	C
ATOM	3611	N	VAL	B	224	61.663	-11.658	7.030	1.00	34.62	N	
ANISOU	3611	N	VAL	B	224	4193	4073	4888	547	351	-582	N
ATOM	3612	CA	VAL	B	224	62.294	-10.430	6.603	1.00	36.49	C	
ANISOU	3612	CA	VAL	B	224	4405	4367	5094	546	438	-647	C
ATOM	3613	C	VAL	B	224	61.405	-9.667	5.848	1.00	40.79	C	
ANISOU	3613	C	VAL	B	224	4995	4969	5534	541	470	-624	C
ATOM	3614	O	VAL	B	224	61.126	-10.130	4.534	1.00	39.49	O	
ANISOU	3614	O	VAL	B	224	4864	4837	5304	575	481	-673	O
ATOM	3615	CB	VAL	B	224	63.659	-10.710	5.962	1.00	38.09	C	
ANISOU	3615	CB	VAL	B	224	4564	4589	5318	594	502	-696	C
ATOM	3616	CG1	VAL	B	224	64.370	-9.405	5.625	1.00	41.85	C	
ANISOU	3616	CG1	VAL	B	224	5008	5121	5773	583	594	-652	C
ATOM	3617	CG2	VAL	B	224	64.511	-11.480	6.961	1.00	41.28	C	
ANISOU	3617	CG2	VAL	B	224	4919	4937	5829	605	461	-715	C
ATOM	3618	N	ALA	B	225	60.994	-8.467	6.062	1.00	37.13	N	
ANISOU	3618	N	ALA	B	225	4535	4521	5053	503	487	-552	N
ATOM	3619	CA	ALA	B	225	60.352	-7.524	5.157	1.00	36.91	C	
ANISOU	3619	CA	ALA	B	225	4544	4551	4931	505	532	-516	C
ATOM	3620	C	ALA	B	225	61.299	-6.847	4.295	1.00	36.49	C	
ANISOU	3620	C	ALA	B	225	4475	4547	4844	523	537	-499	C
ATOM	3621	O	ALA	B	225	62.486	-6.421	4.631	1.00	36.24	O	
ANISOU	3621	O	ALA	B	225	4391	4505	4874	518	681	-497	O
ATOM	3622	CB	ALA	B	225	59.379	-6.649	5.965	1.00	32.80	C	
ANISOU	3622	CB	ALA	B	225	4040	4018	4406	461	502	-446	C
ATOM	3623	N	ALA	B	226	60.757	-6.068	3.231	1.00	38.40	N	
ANISOU	3623	N	ALA	B	226	4758	4845	4987	539	676	-478	N
ATOM	3624	CA	ALA	B	226	61.547	-5.174	2.370	1.00	40.47	C	
ANISOU	3624	CA	ALA	B	226	5012	5157	5208	552	780	-449	C
ATOM	3625	C	ALA	B	226	61.765	-3.826	3.071	1.00	39.95	C	
ANISOU	3625	C	ALA	B	226	4815	4956	5066	506	819	-368	C
ATOM	3626	O	ALA	B	226	61.116	-3.558	4.109	1.00	36.60	O	
ANISOU	3626	O	ALA	B	226	4509	4604	4795	473	762	-336	O

TABLE 3-continued

ATOM	3627	CB	ALA	B	226	60.871	-4.961	0.988	1.00	42.30	C
ANISOU	3627	CB	ALA	B	226	5300	5460	5313	587	809	-446 C
ATOM	3628	N	ARG	B	227	62.705	-3.018	2.530	1.00	34.86	N
ANISOU	3628	N	ARG	B	227	4263	4454	4527	506	916	-339 N
ATOM	3629	CA	ARG	B	227	63.028	-1.824	3.013	1.00	35.97	C
ANISOU	3629	CA	ARG	B	227	4391	4574	4702	461	969	-262 C
ATOM	3630	C	ARG	B	227	63.630	-1.549	4.399	1.00	34.82	C
ANISOU	3630	C	ARG	B	227	4190	4370	4670	417	940	-263 C
ATOM	3631	O	ARG	B	227	62.936	-1.186	5.311	1.00	31.95	O
ANISOU	3631	O	ARG	B	227	3840	3967	4332	387	888	-234 O
ATOM	3632	CB	ARG	B	227	61.807	-0.676	3.043	1.00	38.65	C
ANISOU	3632	CB	ARG	B	227	4535	4655	4734	449	954	-193 C
ATOM	3633	CG	ARG	B	227	61.047	-0.710	1.771	1.00	47.71	C
ANISOU	3633	CG	ARG	B	227	5994	5117	5018	493	967	-185 C
ATOM	3634	CD	ARG	B	227	61.787	-0.069	0.597	1.00	52.83	C
ANISOU	3634	CD	ARG	B	227	6649	6817	6607	511	1075	-155 C
ATOM	3635	NE	ARG	B	227	60.894	0.421	-0.217	1.00	65.59	N
ANISOU	3635	NE	ARG	B	227	8332	8475	8115	539	1073	-108
ATOM	3636	CZ	ARG	B	227	60.252	1.682	-0.276	1.00	65.40	C
ANISOU	3636	CZ	ARG	B	227	8343	8441	8065	528	1108	-20 C
ATOM	3637	NH1	ARG	B	227	60.901	2.714	0.321	1.00	58.59	N
ANISOU	3637	NH1	ARG	B	227	7459	7528	7274	483	1163	34 N
ATOM	3638	NH2	ARG	B	227	59.178	1.895	-1.033	1.00	57.26	N
ANISOU	3638	NH2	ARG	B	227	7369	7455	6931	565	1088	11 N
ATOM	3639	N	PRO	B	228	64.913	01.883	4.527	1.00	25.50	N
ANISOU	3639	N	PRO	B	228	4211	4458	4820	416	974	-299 N
ATOM	3640	CA	PRO	B	228	65.513	-1.761	5.765	1.00	36.34	C
ANISOU	3640	CA	PRO	B	228	42.58	4530	5028	375	951	-300 C
ATOM	3541	C	PRO	B	228	35.522	-0.255	5.241	1.000	35.32	C
ANISOU	3641	C	PRO	B	228	4137	41367	4317	319	988	-228 C
ATOM	3642	O	PRO	B	228	35.331	0.643	5.429	1.00	36.03	O
ANISOU	3642	O	PRO	B	228	4259	4486	4066	315	1036	-183 O
ATOM	3643	CB	PRO	B	228	57.082	-24	12	5.989	1.00	37.72 C
ANISOU	3643	CB	PRO	B	228	43310	4724	52431	391	1012	-341 C
ATOM	3644	CG	PRO	B	228	67.131	-2.204	3.904	1.00	40.	13 C
ANISOU	3644	CG	PRO	B	228	4657	5052	5463	438	1033	-351 C
ATOM	3645	CD	PRO	B	228	55.737	-2.425	3.415	1.00	34.21	C
ANISOU	3645	CD	PRO	B	228	4027	4347	4323	458	1037	-843 C
ATOM	3546	N	GLU	B	229	55.460	-0.022	7.545	1.55	35.35	N
ANISOU	3646	N	GLU	B	229	4121	4324	49913	275	1.00	-220 N

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TABLE 3-continued

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ATOM	3647	CA	GLU	B	229	55.430	1.370	8.090	1.00	32.50	C
ANISOU	3647	CA	GLU	B	229	3753	3031	4655	224	959	-134 C
ATOM	3648	C	GLU	B	229	36.931	1939	8.158	1.00	34.88	C
ANISOU	3648	C	GLU	B	229	3392	4235	5524	185	1042	-165 C
ATOM	3649	O	GLU	B	229	57.383	1.208	8.426	1.00	32.49	O
ANISOU	3649	O	GLU	B	229	47109	4333	5674	151	1.030	-213 O
ATOM	3650	CB	GLU	B	229	64.822	1.345	3.451	1.00	28.86	C
ANISOU	3650	CB	GLU	B	229	3308	3425	4234	197	882	-133 C
ATOM	3651	CG	GLU	B	229	63.029	1.023	9.238	1.00	30.45	C
ANISOU	3651	CG	GLU	B	229	3578	3628	4303	228	328	-150 C
ATOM	3652	CD	GLU	B	229	62.4913	1.133	10.522	1.00	31.92	C
ANISOU	3652	CD	GLU	B	229	3780	3777	45473	201	755	-137 C
ATOM	3653	OE1	GLU	B	229	63.059	11139	11	1618	1.00	37.02 O
ANISOU	3653	OE1	GLU	B	229	4497	4510	5402	153	725	-152 O
ATOM	3654	OE2	GLU	B	229	61.257	1.888	10.379	1.00	34.60	O
ANISOU	3654	OE2	GLU	B	229	4173	4118	4856	216	790	-111 O
ATOM	3655	N	GLY	B	230	67.033	3.256	7.947	1.00	37.21	N
ANISOU	3655	N	GLY	B	230	4175	4389	51	54	145	1114 -111 N
ATOM	3656	CA	GLY	B	230	58.323	9.931	8.284	1.00	35.42	C
ANISOU	3656	CA	GLY	B	230	4134	4405	5298	89	1171	-108 C
ATOM	3657	C	GLY	B	230	58.110	4.564	9.500	.100	135.66	C
ANISOU	3657	C	GLY	B	230	4163	4376	3385	29	139	-88 C
ATOM	3658	O	GLY	B	230	57.005	4.954	15.037	1.00	34.57	O
ANISOU	3658	O	GLY	B	230	3957	4062	51095	1.85	1078	-74 O
ATOM	3659	N	PRO	B	231	69.149	5.587	9.905	1.00	35.25	N
ANISOU	3659	N	PRO	B	231	39130	4132	5280	-31	183	-88 N
ATOM	3660	CA	PRO	B	231	69.062	13	428	11.036	1.00	35.46 C
ANISOU	3660	CA	PRO	B	231	3953	4155	5354	-51	1153	-82 C
ATOM	3661	C	PRO	B	231	37.332	7.491	10.994	1.00	33.75	C
ANISOU	3661	C	PRO	B	231	4209	4270	5486	-100	1168	-24 C
ATOM	3662	O	PRO	B	231	57.457	7.3164	12.046	1.00	35.57	O
ANISOU	3662	O	PRO	B	231	4203	4203	5488	-125	1114	-23 O
ATOM	3663	CB	PRO	B	231	70.406	7.143	11.050	1.00	34.94	C
ANISOU	3663	CB	PRO	B	231	3315	4087	5371	-155	1223	-34 C
ATOM	3664	CG	PRO	B	231	71.347	6.036	10.555	1.00	137.40	C
ANISOU	3664	CG	PRO	B	231	4053	4455	5693	-118	1235	-128 C
ATOM	3665	CD	PRO	B	231	70.538	5.545	9.378	1.00	35.77	C
ANISOU	3665	CD	PRO	B	231	3916	4287	533.7	-4.9	1.757	-106 C
ATOM	3666	N	LEU	B	232	37.535	7.360	9.801	1.00	34.73	N
ANISOU	3666	N	LEU	B	232	4006	4521	5168	-77	1241	23 N

TABLE 3-continued

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ATOM	3667	CA	LEU	B	232	53.623	9.031	9.635	1.00	37.39	C
ANISOU	3667	CA	LEU	B	232	4492	4375	5530	-78	1263	59 C
ATOM	3668	C	LEU	B	232	65.266	8.491	9.142	1.00	36.74	C
ANISOU	3668	C	LEU	B	232	4416	4253	5239	-5	1217	106 C
ATOM	3669	O	LEU	B	232	54.350	9.285	8.918	1.00	34.58	O
ANISOU	3669	O	LEU	B	232	4.214	3951	4977	18	1230	159 O
ATOM	3670	CB	LEU	B	232	67.143	10.059	8.626	1.00	36.74	C
ANISOU	3670	CB	LEU	B	232	4365	4219	5377	-102	1378	150 C
ATOM	3671	CG	LEU	B	232	68.276	10.059	9.144	1.00	44.18	C
ANISOU	3671	CG	LEU	B	232	5252	5117	6416	-189	1431	147 C
ATOM	3672	CD1	LEU	B	232	68.341	11.857	8.028	1.00	41.32	C
ANISOU	3672	CD1	LEU	B	232	4906	4749	6.044	-215	1554	211 C
ATOM	3673	CD2	LEU	B	232	67.876	11.733	10.393	1.00	39.17	C
ANISOU	3673	CD2	LEU	B	232	4637	4408	5838	-234	1384	139 C
ATOM	3674	N	THR	B	233	65.142	7.170	3.955	1.00	35.57	N
ANISOU	3674	N	THR	B	233	4262	470	522	41	1164	62 N
ATOM	3675	CA	THR	B	233	63.916	6.637	8.377	1.00	34.58	C
ANISOU	3675	CA	THR	B	233	4184	4061	4895	104	1124	74 C
ATOM	3676	C	THR	B	233	83.320	5.542	41.260	1.00	32.04	C
ANISOU	3676	C	THR	B	233	3849	3743	4580	122	1017	23 C
ATOM	3677	O	THR	B	233	6.2.577	4.794	3.791	1.00	32.13	O
ANISOU	3677	O	THR	B	233	3887	3789	4531	170	976	9 O
ATOM	3678	CB	THR	B	233	64.154	6.072	6.001	1.00	40.97	C
ANISOU	3678	CB	THR	B	233	5000	4934	55141	151	1175	75 C
ATOM	3679	OG1	THR	B	233	65.209	5.104	6.999	1.00	38.11	O
ANISOU	3679	OG1	THR	B	233	4567	4602	5311	150	1173	14 O
ATOM	3680	CG2	THR	B	233	64.550	7.165	5.897	1.00	40.19	C
ANISOU	3680	CG2	THR	B	233	4929	4838	5502	141	1286	141 C
ATOM	3681	N	GLY	B	234	63.553	5.578	10.660	1.00	39.81	N
ANISOU	3681	N	GLY	B	234	3659	35233	4496	82	069	-3 N
ATOM	3682	CA	GLY	B	234	52.904	4.603	11.414	1.00	29.36	C
ANISOU	3682	OA	GLY	B	234	3470	3371	4313	97	871	-38 O
ATOM	3683	C	GLY	B	234	61.408	4.857	11.380	1.00	29.81	C
ANISOU	3683	C	GLY	B	234	3531	33601	4245	124	838	-15 C
ATOM	3684	O	GLY	B	234	60.992	5.017	11.292	1.00	30.66	O
ANISOU	3684	O	GLY	B	234	3742	3504	4404	114	877	41 O
ATOM	3685	N	GLY	B	234	63.597	3.832	11.443	1.00	28.35	N
ANISOU	3685	N	GLY	B	235	3423	3263	4089	157	763	-28 N
ATOM	3686	CA	GLY	B	235	59.134	3.994	11.467	1.00	26.70	C
ANISOU	3686	CA	GLY	B	235	3266	3057	3823	180	731	0 C

TABLE 3-continued

ATOM	3687	C	GLY	B	235	58.509	4.311	10.149	1.00	31.63	C
ANISOU	3687	C	GLY	B	235	3939	3712	4366	223	771	37 C
ATOM	3688	O	GLY	B	235	57.313	4.501	10.090	1.00	32.59	O
ANISOU	3688	O	GLY	B	235	4100	3843	4440	247	742	62 O
ATOM	3689	N	ALA	B	236	59.305	4.341	9.065	1.00	21.92	N
ANISOU	3689	N	ALA	B	236	3602	3402	4012	236	837	41 N
ATOM	3690	CA	ALA	B	236	59.777	4.761	7.737	1.00	31.11	C
ANISOU	3690	CA	ALA	B	236	3921	3707	4164	276	884	84 C
ATOM	3691	C	ALA	B	236	57.649	0.319	7.416	1.00	32.51	C
ANISOU	3691	C	ALA	B	236	4120	3925	4306	1319	813	63 C
ATOM	3692	O	ALA	B	236	57.775	2.602	7.666	1.00	33.11	O
ANISOU	3692	O	ALA	B	236	4188	4314	4399	320	760	5 O
ATOM	3693	CB	ALA	B	236	59.876	4.559	8.660	1.00	37.54	C
ANISOU	3693	CB	ALA	B	236	4717	4554	4991	287	956	75 C
ATOM	3694	N	PHE	B	237	56.581	4.524	6.790	1.90	55.24	N
ANISOU	3694	N	PHE	B	237	4516	4296	4579	354	814	107 N
ATOM	3695	CA	PHE	B	237	55.422	3.486	6.583	1.00	52.65	C
ANISOU	3695	CA	PHE	B	237	4203	4010	4154	366	741	65 C
ATOM	3596	C	PHE	B	237	55.803	2.351	5.577	1.00	34.92	C
ANISOU	3696	C	PHE	B	237	4484	4348	4437	4121	733	34 C
ATOM	3697	O	PHE	B	237	55.388	1.213	5.751	1.30	33.9	O
ANISOU	3697	O	PHE	B	237	4343	4234	4306	416	670	-18 O
ATOM	3698	CB	PHE	B	237	54.272	4.355	6.046	1.09	35.59	C
ANISOU	3698	CB	PHE	B	237	4524	4403	4494	42.3	74.6	145 C
ATOM	3699	CG	PHE	B	237	53.021	35.70	5.741	1.00	34.15	C
ANISOU	3699	CG	PHE	B	237	4453	4274	4249	455	571	126 C
ATOM	3700	CD1	PHE	B	237	52.527	2.625	6.662	1.33	33.86	C
ANISOU	3700	CD1	PHE	B	237	4390	4232	4252	432	592	7.3 C
ATOM	3701	CD2	PHE	B	237	52.339	2.745	4.500	1.00	36.10	C
ANISOU	3701	CD2	PHE	B	237	4739	4583	4393	506	679	154 C
ATOM	3702	CE1	PHE	B	237	51.374	1.868	6.380	1.00	30.19	C
ANISOU	3702	CE1	PHE	B	237	4687	4573	4492	40	523	57 C
ATOM	3703	CE2	PHE	B	237	51.150	3.009	4.239	1.00	33.60	C
ANISOU	3703	CE2	PHE	B	237	4.425	4320	4023	531	654	129 C
ATOM	3704	CZ	PHE	B	237	50.689	2.055	5.1	65.	1.00	3.534 C
ANISOU	3704	CZ	PHE	B	237	4612	4527	4288	501	527	79 C
ATOM	3705	N	ALA	B	238	50.637	2.553	4.523	1.00	35.58	N
ANISOU	3705	N	ALA	B	238	4701	4551	4516	426	815	47 N
ATOM	3706	CA	ALA	B	238	57.146	1.651	3.633	1.00	35.45	C
ANISOU	3706	CA	ALA	B	238	4550	4437	4432	453	824	-7 C

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TABLE 3-continued

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ATOM	3707	C	ALA	B	238	57.883	9.459	4.331	1.00	3.53	C	
ANISOU	3707	C	ALA	B	238	5023	4385	5033	433	764	-62	C
ATOM	3708	O	ALA	B	238	57.765	-0.711	3.916	1.00	41.00	O	
ANISOU	3708	O	ALA	B	238	5200	5191	5188	456	744	-143	O
ATOM	3709	CB	ALA	B	238	58.044	2.353	2.5130	1.00	42.20	C	
ANISOU	3709	CB	ALA	B	238	5415	5367	52513	454	927	28	C
ATOM	3710	N	SER	B	239	58.595	0.722	5.435	1.00	33.60	N	
ANISOU	3710	N	SER	B	239	4225	4173	4359	393	768	-60	N
ATOM	3711	CA	SER	B	239	59.248	-0.331	6.200	1.00	32.29	C	
ANISOU	3711	CA	SER	B	239	4010	3951	4277	375	744	-144	C
ATOM	3712	C	SER	B	239	58.205	-1.033	7.073	1.03	31.75	C	
ANISOU	3712	C	SER	B	239	354	3857	422.3	365	646	-159	C
ATOM	3713	O	SER	B	239	58.241	-2.234	7.228	1.00	32.74	O	
ANISOU	3713	O	SER	B	239	4059	4012	4357	373	594	-213	O
ATOM	3714	CB	SER	B	239	60.373	0.225	7.0136	1.30	32.38	C	
ANISOU	3714	CB	SER	B	239	3974	3953	4377	334	7313	-132	C
ATOM	3715	OG	SER	B	239	61.340	0.328	6.173	1.00	32.63	O	
ANISOU	3715	OG	SER	B	239	3995	4037	4395	337	374	-115	O
ATOM	3716	N	LYS	B	240	57.267	-0.295	7.640	1.00	31.06	N	
ANISOU	3716	N	LYS	B	240	3983	3791	4126	350	624	-114	N
ATOM	3717	CA	LYS	B	240	56.362	-0.946	8.020	1.00	33.63	C	
ANISOU	3717	CA	LYS	B	240	4212	4104	4431	334	536	025	C
ATOM	3718	C	LYS	B	240	55.337	-1.803	7.857	1.130	32.36	C	
ANISOU	3718	C	LYS	B	240	4073	3975	4245	3713	485	-154	C
ATOM	3719	O	LYS	B	240	54.739	-2.806	8.430	1.00	31.45	O	
ANISOU	3719	O	LYS	B	240	3343	3842	4144	3.51	414	-167	O
ATOM	3720	CB	LYS	B	240	55.754	0.069	9.519	1.00	29.45	C	
ANISOU	3720	CB	LYS	B	240	3635	3545	3959	310	529	-79	C
ATOM	3721	CG	LYS	B	240	55.554	0.913	10.488	1.50	29.47	C	
ANISOU	3721	CG	LYS	B	240	3661	3502	4033	272	561	-61	C
ATOM	3722	CD	LYS	B	240	57.407	3.112	11.4617	1.00	32.01	C	
ANISOU	3722	CD	LYS	B	240	3934	3794	4429	244	525	-102	C
ATOM	3723	CE	LYS	B	240	58.753	-0.202	13.831	1.03	12.35	C	
ANISOU	3723	CE	LYS	B	240	4106	4003	4655	253	573	-131	C
ATOM	3724	NZ	LYS	B	240	59.557	-0.890	11.372	1.00	32.00	N	
ANISOU	3724	NZ	LYS	B	240	3854	3773	4530	230	532	-103	N
ATOM	3725	N	VAL	B	241	54.535	-1.373	5.694	1.00	32.42	N	
ANISOU	3725	N	VAL	B	241	4114	4032	4172	397	522	-136	N
ATOM	3726	CA	VAL	B	241	53.884	-2.062	5.916	1.00	33.51	C	
ANISOU	3726	CA	VAL	B	241	4289	4228	4247	424	474	-161	C

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TABLE 3-continued

ATOM	3727	C	VAL	B	241	54.309	-3.473	5.520	1.00	C
ANISOU	3727	C	VAL	B	241	4684	46.36	4660	433	444
						-238	C			
ATOM	3728	O	VAL	B	241	53.509	-4.42.6	5.602	3.00	133.30
ANISOU	3728	O	VAL	B	241	4247	4187	4210	428	3713
						-276	O			
ATOM	3729	CB	VAL	B	241	53.357	-1.214	4.690	1.00	35.23
ANISOU	3729	CB	VAL	B	241	4536	4495	435.5	4515	516
						-118	C			
ATOM	3730	CG1	VAL	B	241	54.260	-1.2190	3.427	1.00	35.39
ANISOU	3730	CG1	VAL	B	241	4568	4553	4324	497	582
						-135	C			
ATOM	3731	CG2	VAL	B	241	51.976	-1.674	4.308	1.00	31.73
ANISOU	3731	CG3	VAL	B	241	5252	52318	4981	482	448
						-130	C			
ATOM	3732	N	GLY	B	242	50590	-3.621	5.177	1.00	21.815
ANISOU	3732	N	GLY	B	242	4041	4.013	4059	443	498
						-263	N			
ATOM	3733	CA	GLY	B	242	55.070	-4.980	4.919	1.00	24.25
ANISOU	3733	CA	GLY	B	242	43123	4236	41384	456	470
						-341	C			
ATOM	3734	C	GLY	B	242	56.056	-5.044	5	80	1.00
ANISOU	3734	C	GLY	B	242	4022	3952	4185	425	402
						-365	C			
ATOM	3735	O	GLY	B	242	55.750	-7.033	6.045	1.00	35.76
ANISOU	3735	O	GLY	B	242	4504	4416	4566	400	346
						-220.	O			
ATOM	3736	N	ALA	B	243	56.428	-5.258	7.313	1.00	29.33
ANISOU	3736	N	ALA	B	243	3656	3574	3914	392	405
						-225	N			
ATOM	3737	CA	ALA	B	243	56.430	-5.025	8.554	1.00	3.29
ANISOU	3737	CA	ALA	B	243	3754	3643	4112	353	341
						-338	C			
ATOM	3738	C	ALA	B	243	55.025	-6.444	8.914	1.00	30.49
ANISOU	3738	C	ALA	B	243	3802	3558	4116	344	259
						-333	C			
ATOM	3739	O	ALA	B	243	54.807	-7.598	9.334	1.00	32.27
ANISOU	3739	O	ALA	B	243	4024	3851	4877	332	208
						-368	O			
ATOM	3740	CB	ALA	B	243	57.033	-5.11-17	9.712	1.00	31.31
ANISOU	3740	CB	ALA	B	243	3852	3741	4304	331	359
						-296	C			
ATOM	3741	N	LEU	B	244	54.0148	105	-5.526	8.784	1.00
ANISOU	3741	N	LEU	B	244	3864	37.45	4105	340	276
						-287	N			
ATOM	3742	CA	LEU	B	244	52.674	-5.909	8.960	1.00	30.99
ANISOU	3742	CA	LEU	B	244	3398	3779	4096	325	212
						-284	C			
ATOM	3743	C	LEU	B	244	52.239	-5.955	7.957	1.00	29.67
ANISOU	3743	C	LEU	B	244	3751	3636	3886	344	178
						-041	C			
ATOM	3744	O	LEU	B	244	51.449	-7.820	8.324	1.00	32.72
ANISOU	3744	O	LEU	B	244	4136	4009	4286	320	112
						-363	O			
ATOM	3745	CB	LEU	B	244	51.744	-4.689	8.338	1.00	28.42
ANISOU	3745	CB	LEU	B	244	358.8	3494	3715	3.31	230
						-226	C			
ATOM	3746	CG	LEU	B	244	51.973	-3.742	10.052	1.00	20.20
ANISOU	3746	CG	LEU	B	244	3759	3649	3954	305	249
						-175	C			

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ATOM	3747	CD1	LEU	B	244	51.316	-2.399	9.689	1.07	30.96	C	
ANISOU	3747	CD1	LEU	B	244	3910	3617	4033	324	287	-121 C	
ATOM	3748	CD2	LEU	B	244	51.466	-4.153	11.421	1.00	26.79	C	
ANISOU	3748	CD2	LEU	B	244	3345	3225	3609	264	192	-169 C	
ATOM	3749	N	LEU	B	245	52.662	-6.860	6.712	1.00	30.85	N	
ANISOU	3749	N	LEU	B	245	3918	3824	3980	383	222	-355 N	
ATOM	3750	CA	LEU	B	245	52.347	-7.946	5.789	1.00	32.43	C	
ANISOU	3750	CA	LEU	B	245	4137	4044	4141	400	183	-435 C	
ATOM	3751	C	LEU	B	245	52.875	9.325	6.287	1.00	34.89	C	
ANISOU	3751	C	LEU	B	245	4435	4289	4530	386.	145	-495 C	
ATOM	3752	O	LEU	B	245	52.130	-10.307	5.195	1.00	33.79	O	
ANISOU	3752	O	LEU	B	245	4310	4140	4389	371	83	-538 O	
ATOM	3753	CB	LEU	B	245	52.861	-7.564	4.345	1.00	33.36	C	
ANISOU	3753	CB	LEU	B	245	4277	4219	4161	449	249	-458 C	
ATOM	3754	CG	LEU	B	245	52.626	-8.836	3.291	1.00	36.13	C	
ANISOU	3754	CG	LEU	B	245	4048	4568	4490	458	213	-642 C	
ATOM	3755	CD1	LEU	B	245	51.096	-9.0981	3.109	1.00	35.71	C	
ANISOU	3755	CD1	LEU	B	245	4513	4-576	4379	454	143	-551 C	
ATOM	3756	CD2	LEU	B	245	513.059	-8.553	2.101	1.00	37.95	C	
ANISOU	3756	CD2	LEU	B	245	4897	4874	4651	518	283	-565 C	
ATOM	3757	N	LEU	B	246	54.103	-9.416	13.834	1.00	34.18	N	
ANISOU	3757	NN	LEU	B	246	4322	4154	4511	390i	175	-407 N	
ATOM	3758	CA	LEU	B	246	54.594	-10.677	7.409	1.00	35.73	C	
ANISOU	3758	CA	LEU	B	246	4505	4231	4787	382	130	-342 C	
ATOM	3759	C	LEU	B	246	53.587	-11.247	8.480	1.00	06.74	C	
ANISOU	3759	C	LEU	B	246	4635	4370	4953	333	52	-524 C	
ATOM	3760	O	LEU	B	246	53332	-12.456	53.515	1.00	35.03	O	
ANISOU	3760	O	LEU	B	246	4431	4112	4785	324	-1	-571 O	
ATOM	3761	CB	LEU	B	246	55.955	-10.467	8.112	1.00	38.46	C	
ANISOU	3761	CB	LEU	B	246	481.6	4592	15205	383	618	-528 C	
ATOM	3762	CG	LEU	B	246	57.082	-9.983	7.158	1.00	43.58	C	
ANISOU	3762	CG	LEU	B	246	5463	5287	5636	432	251	-547 C	
ATOM	3763	CD	LEU	B	246	1	58.437	-9.871	7.343	1.00	47.33	C
ANISOU	3763	CD1	LEU	B	246	5843	5882	6344	435	281	-540 C	
ATOM	3764	CD2	LEU	B	246	57.200	-10.866	5.936	1.00	42.81	C	
ANISOU	3764	CD2	LEU	B	246	5379	5138	5593	474	256	-624 C	
ATOM	3765	N	VAL	B	247	53.032	-10.355	9.330	1.00	34.15	N	
ANISOU	3765	N	VAL	B	247	4200	4046	4518	392	49	-467 N	
ATOM	3766	CA	VAL	B	247	52.075	-10.770	10.343	1.00	31.36	C	
ANISOU	3766	CA	VAL	B	247	3940	367.5	4301	255	-15	-431 C	

TABLE 3-continued

ATOM	3767	C	VAL	B	247	50.797	-11.311	9.709	1.00	31.79	C		
ANISOU	3767	C	VAL	B	247	4015	3757	4306	241	-61	-459	C	
ATOM	3768	O	VAL	B	247	50.335	-12424	10.080	3.00	31.15	C		
ANISOU	3768	O	VAL	B	247	3946	3634	4254	210	-120	-484	O	
ATOM	3769	CB	VAL	B	247	51.750	-9.578	11.30	1.00	30.29	C		
ANISOU	3769	CB	VAL	B	247	3787	3557	4155	232	2	-358	C	
ATOM	3770	CG1	VAL	B	247	50.573	-9.587	12.192	1	00	29.85	C	
ANISOU	3770	CG1	VAL	B	247	3723	3495	4119	186	-55	-331	C	
ATOM	3771	CG2	VAL	B	247	52.953	-9.158	12.113	1.00	30.60	C		
ANISOU	3771	CG2	VAL	B	247	3552	3554	4252	234	34	-07.5	C	
ATOM	3772	N	ASP	B	248	50.240	-10.556	15.741	1.00	32.08	N		
ANISOU	3772	N	ASP	B	248	4064	3865	4260	264	-351	-454	N	
ATOM	3773	CA	ASP	B	248	49.096	-10.952	6.022	1.60	34.71	C		
ANISOU	3773	CA	ASP	B	248	4411	4239	4537	256	-77	-483	C	
ATOM	3774	C	ASP	B	248	48.264	-12.330	7	371	1.00	36.102	C	
ANISOU	3774	C	ASP	B	248	4635	4411	4754	259	-113	-568	C	
ATOM	3775	O	ASP	B	248	48.379	-13.215	7.521	1.00	33.42	O		
ANISOU	3775	O	ASP	B	248	4270	4027	4431	220	-177	-595	O	
ATOM	3776	CB	ASP	B	248	48.739	-9.880	6.979	1.00	35.25	C		
ANISOU	3776	CB	ASP	B	248	4493	4394	4511	294	-38	-464	C	
ATOM	3777	CG	ASP	B	248	47.369	-10.155	6.359	1.00	35.59	C		
ANISOU	3777	CG	ASP	B	248	4657	4617	4617	280	1.00	-454	C	
ATOM	3778	OD1	ASP	B	248	45.411	-1	0.41	8	7.124	1.00	35.43	O
ANISOU	3778	OD1	ASP	B	248	4525	4567	4518	239	-140	-465	O	
ATOM	3779	OD2	ASP	B	248	47.259	-10.096	5.1137	1.00	40.24	O		
ANISOU	3779	OD2	ASP	B	248	5150	5.137	5004	319	-79	-518	O	
ATOM	3780	N	VAL	B	249	50.354	-12.527	6.622	1.00	03.37	N		
ANISOU	3780	N	VAL	B	249	4273	4031	4376	303	-71	-613	N	
ATOM	3781	CA	VAL	B	249	50.559	-1.3	8.831	5.052	1.00	33.10	C	
ANISOU	3781	CA	VAL	B	249	4548	4351	4741	313	-132	-703	C	
ATOM	3782	C	VAL	B	249	50.651	-14.033	5.952	1.00	38.79	C		
ANISOU	3782	C	VAL	B	249	4725	4334	4921	276	-355	-716	C	
ATOM	3783	O	VAL	B	249	50.132	-16.121	6.762	1.00	37.04	O		
ANISOU	3783	O	VAL	B	249	4775	4328	4971	263	-231	-775	O	
ATOM	3784	CB	VAL	B	249	51.833	-13.700	5.066	1.00	37.92	C		
ANISOU	3784	CB	VAL	B	249	41377	4537	4542	374	-34	-742	C	
ATOM	3785	CG1	VAL	B	249	52.289	-15.334	4.545	1.00	45.49	C		
ANISOU	3785	CG1	VAL	B	249	5856	5498	5920	393	-57	-837	C	
ATOM	3786	CG2	VAL	B	249	51.480	-12.712	3.890	1.00	37.75	C		
ANISOU	3786	CG2	VAL	B	249	4871	4664	4307	405	6	-736	C	

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ATOM	3787	N	LEU	B	250	51.288	-14.635	8.079	100.	32.20	N
ANISOU	3787	N	LEU	B	250	4120	3712	4404	268	-141	-662 N
ATOM	3788	CA	LEU	B	250	51.551	-15.753	9.037	1.00	35.50	C
ANISOU	3788	CA	LEU	B	250	45136	40.38	4913	243	-185	-169 C
ATOM	3789	C	LEU	B	250	50.198	6.177	9.503	1.00	35.32	C
ANISOU	3789	C	LEU	B	250	4518	4007	4894	161	-249	-651 C
ATOM	3790	O	LEU	B	250	49.911	-17.334	9.532	1.00	34.44	O
ANISOU	3790	O	LEU	B	250	4424	3837	4823	1.57	-298	-692 O
ATOM	3791	CB	LEU	B	250	52.422	-15.281	13.162	3.00	34.53	C
ANISOU	3791	CB	LEU	B	250	4398	3898	4832	244	-162	-608 C
ATOM	3792	CG	LEU	B	250	52.445	-13.1198	11.409	1.00	35.16	C
ANISOU	3792	CG	LEU	B	250	4462	3884	5015	210	-216	-584 C
ATOM	3793	CD1	LEU	B	250	53.006	-17.581	11.155	1.00	33.41	C
ANISOU	3793	CD1	LEU	B	250	426.1	3580	4855	230	-245	-550 C
ATOM	3794	CD2	LEU	B	250	53.231	-15.473	12.525	1.00	34.79	C
ANISOU	3794	CD2	LEU	B	250	4382	3829	5006	211	-193	-520 C
ATOM	3795	N	VAL	B	251	49.353	-15.238	9.906	1.000	0033.71	N
ANISOU	3795	N	VAL	B	251	4297	3861	4652	154	-245	-589 N
ATOM	3796	CA	VAL	B	251	48.130	-15.655	10.595	1.00	34.72	C
ANISOU	3796	CA	VAL	B	251	4419	3979	4795	90	-303	-553 C
ATOM	3797	C	VAL	B	251	47.117	-16.223	9.544	1.00	35.45	C
ANISOU	3797	C	VAL	B	251	4527	4104	4339	74	-342	-525 C
ATOM	3798	O	VAL	B	251	46.390	-17.179	9.809	1.00	37.83	O
ANISOU	3798	O	VAL	B	251	4839	4372	5182	22	-398	-544 O
ATOM	3799	CB	VAL	B	251	47.570	-14.486	11.496	1.00	33.03	C
ANISOU	3799	CB	VAL	B	251	4175	13814	4551	69	-287	-477 C
ATOM	3800	CG1	VAL	B	251	46.165	-14.827	12.035	1.00	28.03	C
ANISOU	3800	CG1	VAL	B	251	3529	3193	3928	6	-3.39	-454 C
ATOM	3801	CG2	VAL	B	251	48.641	-14.129	12.512	1.00	32.54	C
ANISOU	3801	CG2	VAL	B	251	4101	3711	4551	81	-258	-433 C
ATOM	3802	N	ASN	B	252	47.061	-15.633	0.340	1.00	28.19	N
ANISOU	3802	N	ASN	B	252	4832	4526	5104	116	-3.14	-656 N
ATOM	3803	CA	ASN	B	252	46.256	-16.231	7.272	1.00	37.04	C
ANISOU	3803	CA	ASN	B	252	475.2	4413	4900	107	-353	-727 C
ATOM	3804	C	ASN	B	252	46.755	-17	621	6.885	1.00	38.88 C
ANISOU	3804	C	ASN	B	252	5016	4567	5191	105	-332	-314 C
ATOM	3805	O	ASN	B	252	45.930	-18.434	6.542	1.00	37.12	O
ANISOU	3805	O	ASN	B	252	4803	4332	4969	65	-438	-857 O
ATOM	3806	CB	ASN	B	252	415.254	-15.3.70	6.019	1.00	38.03	C
ANISOU	3806	CB	ASN	B	252	4085	4635	4933	1150	-315	-744 C

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ATOM	3807	CG ASN B	252	45.683	-13.962	6.283	1.00	39.82	C
ANISOU	3807	CG ASN B	252	5057	49361	5105	158	-287	-659
ATOM	3808	OD1 ASN B	252	45.053	-13.730	7.322	1.00	35.44	O
ANISOU	3808	OD1 ASN B	252	46133	4502	4711	126	-307	-502
ATOM	3809	ND2 ASN B	252	46.030	-1	3.08	5.4133	1.00	34.41
ANISOU	3809	ND2 ASN B	252	46.37	4572	4597	223	-235	-548
ATOM	3810	N SER B	253	48.073	-17.854	6.902	1.00	41.14	N
ANISOU	3810	N SER B	253	5314	4799	5513	151	-345	-333
ATOM	3811	CA SER B	253	48.543	-19	281	5.675	1.00	44.01
ANISOU	3811	CA SER B	253	57081	5052	504.5	152	-376	-916
ATOM	3812	C SER B	253	48.138	-20.212	7.797	1.00	41.84	C
ANISOU	3812	C SER B	253	5434	4733	5764	44	-432	-892
ATOM	3813	O SER B	253	47.760	-21.1428	7.524	1.00	42.34	O
ANISOU	3813	O SER B	253	5522	4706	58.59	61	-480	-955
ATOM	3814	CB SER B	253	50.336	-19.406	6.432	1.00	40.42	C
ANISOU	3814	CB SER B	253	5262	4578	55.19	220	-326	-946
ATOM	3815	OG SER B	253	50.4213	-18.469	5.457	1.30	46.63	O
ANISOU	3815	OG SER B	253	5645	54.54	5217	273	-206	-956
ATOM	3816	N LEU B	254	48.189	-19.746	9.049	1.00	39422	N
ANISOU	3816	N LEU B	254	5076	4353	5472	66	-426	-809
ATOM	3817	CA LEU B	254	47.536	-20.533	10.168	1.00	67.62	C
ANISOU	3817	CA LEU B	254	4873	4075	5345	-1	-473	-752
ATOM	3818	C LEU B	254	46.175	-20.833	9.027	1.00	38197	C
ANISOU	3818	C LEU B	254	5041	4277	5483	-68	-527	-777
ATOM	3819	O LEU B	254	45.736	-22.014	10.067	1.140	38.013	O
ANISOU	3819	O LEU B	254	42	4382	5.25	-119	-579	-310
ATOM	3820	CB LEU B	254	47.745	-19.786	11.484	1.00	35.92	C
ANISOU	3820	CB LEU B	254	4628	3868	5151	-16	-461	-659
ATOM	3821	CG LEU B	254	49.094	-191935	12.204	100	36.45	C
ANISOU	3821	CG LEU B	254	4695	3677	5282	24	-438	-633
ATOM	3822	CD1 LEU B	254	49.298	-16.742	13.107	1.99	39.13	C
ANISOU	3822	CD1 LEU B	254	5600	4257	5509	23	-405	-545
ATOM	3823	CD2 LEU B	254	49.177	-2112913	12.378	1.00	36.80	C
ANISOU	3823	CD2 LEU B	254	4764	3799	5420	-7	-480	-837
ATOM	3824	N LEU B	255	451408	-19.702	9.558	1.00	35.65	N
ANISOU	3824	N LEU B	255	4974	4347	5355	-6.0	-512	-753
ATOM	3825	CA LEU B	255	44.975	-20.010	9.251	1.06	41.30	C
ANISOU	3825	CA LEU B	255	5296	4729	5668	-122	-561	-770
ATOM	3826	C LEU B	255	43.734	-21.028	8.003	1.00	44.33	C
ANISOU	3826	C LEU B	255	5774	5105	51135	-137	-601	-882

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ATOM	3827	O	LEU	B	255	42.789	-21.635	8.108	1.00	42	25	O
ANISOU	3827	O	LEU	B	255	5442	4806	5804	-2211	-557	-911	O
ATOM	3828	CB	LEU	B	255	43.204	-18.673	3.984	1.00	39.52	C	
ANISOU	3828	CB	LEU	B	255	5036	4528	5351	-114	-537	-726	C
ATOM	3829	CG	LEU	B	255	42.958	-17.805	10.223	1.90	42.01	C	
ANISOU	3829	CG	LEU	B	255	5316	4063	5678	-134	-518	-623	C
ATOM	3830	CD1	LEU	B	255	42.744	-11.334	94514	1.00	43.432	C	
ANISOU	3830	CD1	LEU	B	255	5524	5306	5821	-86	-475	-566	C
ATOM	3831	CD2	LEU	B	255	41.745	-18.372	16.930	1.00	30.18	C	
ANISOU	3831	CD2	LEU	B	255	4931	4501	5355	-220	-553	-595	C
ATOM	3332	N	GLU	B	256	44.593	-254991	7.083	1.00	45.23	N	
ANISOU	3832	N	GLU	B	256	5352	5216	6118	-58	-570	-947	N
ATOM	3833	CA	GLU	B	256	44.479	-21.926	5.968	1.06	47.83	C	
ANISOU	3833	CA	GLU	B	256	6214	5527	0432	-57	-6135	-1001	C
ATOM	3834	C	GLU	B	256	44.841	-23.360	5.361	1.00	47.22	C	
ANISOU	3834	C	GLU	B	256	6171	5339	5462	-97	-630	-1107	C
ATOM	3835	O	GLU	B	256	44.123	-24.284	6.030	1.00	51.41	O	
ANISOU	3835	O	GLU	B	256	6717	5803	7013	-120	-694	-1172	O
ATOM	3836	CB	GLU	B	256	45.318	-21.435	4.790	1.00	48.10	C	
ANISOU	3836	CB	GLU	B	256	6268	5610	6390	10	-554	-1114	C
ATOM	3837	CG	GLU	B	256	45.167	-22.219	3.508	1.00	61.20	C	
ANISOU	3837	CG	GLU	B	256	7950	7287	8035	23	-5313	-1238	C
ATOM	3838	CD	GLU	B	256	46.023	-21.556	2.412	1.00	613.02	C	
ANISOU	3838	CD	GLU	B	256	8958	8344	8913	117	-524	-1279	C
ATOM	3839	OE1	GLU	B	256	47.209	-224355	2.328	1.00	77.05	O	
ANISOU	3839	OE1	GLU	B	256	0007	9298	9971	508	-487	-1315	O
ATOM	3840	OE2	GLU	B	256	45.563	-20.759	1.563	1.00	75.93	O	
ANISOU	3840	OE2	GLU	B	256	9054	94744	9813	139	-509	-1270	O
ATOM	3841	N	SER	B	257	45.044	-23.560	7.059	1.00	45.90	N	
ANISOU	3841	N	SER	B	257	6015	5059	6306	-53	-611	-1075	N
ATOM	3842	CA	SER	B	257	46.300	-24.912	7.487	1.00	48.42	C	
ANISOU	3842	CA	SER	B	257	6369	5237	6790	-84	-646	-1110	C
ATOM	3843	C	SER	B	257	45.432	-25.477	8.509	1.00	49.77	C	
ANISOU	3843	C	SER	B	257	6532	5345	7034	-177	-694	-1048	C
ATOM	3844	O	SER	B	257	45.288	-26.680	8.707	1.00	47.14	O	
ANISOU	3844	O	SER	B	257	6231	4904	6775	-217	-737	-1090	O
ATOM	3845	CB	SER	B	257	47.764	-25.004	7.913	1.00	47.28	C	
ANISOU	3845	CB	SER	B	257	6235	5026	6703	-14	-606	-1092	C
ATOM	3846	OG	SER	B	257	48.622	-24.230	7.093	1.00	44.81	O	
ANISOU	3846	OG	SER	B	257	5916	4786	6232	68	-546	-1121	O

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TABLE 3-continued

ATOM	3847	N	TYR	B	258	44.874	-24.615	9.456	1.00	44.35	N
ANISOU	3847	N	TYR	B	258	5803	4722	6325	-210	-685	-948
ATOM	3848	CA	TYR	B	258	44.055	-25.037	10.587	1.00	42.12	C
ANISOU	3848	CA	TYR	B	258	5507	4395	6102	-297	-721	-878
ATOM	3849	C	TYR	B	258	42.753	-24.292	10.674	1.00	46.14	C
ANISOU	3849	C	TYR	B	258	5969	5014	6549	-352	-732	-837
ATOM	3850	O	TYR	B	258	42.578	-23.379	11.539	1.00	43.90	O
ANISOU	3850	O	TYR	B	258	5648	4786	6245	-355	-705	-743
ATOM	3851	CB	TYR	B	258	44.820	-24.899	11.877	1.00	42.85	C
ANISOU	3851	CB	TYR	B	258	5596	4433	6252	-282	-699	-784
ATOM	3852	CG	TYR	B	258	46.081	-25.680	11.875	1.00	45.24	C
ANISOU	3852	CG	TYR	B	258	5939	4527	6625	-226	-694	-818
ATOM	3853	CD1	TYR	B	258	46.044	-27.069	12.139	1.00	49.80	C
ANISOU	3853	CD1	TYR	B	258	6557	5071	7295	-265	-739	-843
ATOM	3854	CD2	TYR	B	258	47.310	-25.082	11.593	1.00	43.27	C
ANISOU	3854	CD2	TYR	B	258	5886	4400	6353	-135	-644	-828
ATOM	3855	CE1	TYR	B	258	47.211	-27.834	12.124	1.00	47.89	C
ANISOU	3855	CE1	TYR	B	258	8352	4721	7122	-205	-738	-877
ATOM	3856	CE2	TYR	B	258	48.488	-25.837	11.585	1.00	48.97	C
ANISOU	3856	CE2	TYR	B	258	6439	5025	7143	-78	-640	-863
ATOM	3857	CZ	TYR	B	258	48.401	-27.216	11.824	1.00	49.72	C
ANISOU	3857	CZ	TYR	B	258	6575	4989	7328	-111	-687	-890
ATOM	3858	OH	TYR	B	258	49.499	-27.985	11.819	1.00	53.37	O
ANISOU	3858	OH	TYR	B	258	7067	5351	7861	-50	-686	-924
ATOM	3859	N	PRO	B	259	41.807	-24.681	9.792	1.00	45.61	N
ANISOU	3859	N	PRO	B	259	5899	4981	6450	-394	-772	-910
ATOM	3860	CA	PRO	B	259	40.567	-23.968	9.670	1.00	46.11	C
ANISOU	3860	CA	PRO	B	259	5911	5161	6446	-435	-785	-884
ATOM	3861	C	PRO	B	259	39.792	-23.826	10.991	1.00	44.68	C
ANISOU	3861	C	PRO	B	259	5691	4984	6302	-507	-792	-782
ATOM	3862	O	PRO	B	259	39.013	-22.895	11.113	1.00	43.30	O
ANISOU	3862	O	PRO	B	259	5457	4917	6067	-515	-782	-737
ATOM	3863	CB	PRO	B	259	39.791	-24.801	8.621	1.00	47.63	C
ANISOU	3863	CB	PRO	B	259	6115	5356	5627	-481	-840	-991
ATOM	3864	CG	PRO	B	259	40.832	-25.414	7.784	1.00	48.18	C
ANISOU	3864	CG	PRO	B	259	6242	5358	6708	-421	-834	-1084
ATOM	3865	CD	PRO	B	259	41.934	-25.742	8.768	1.00	49.33	C
ANISOU	3865	CD	PRO	B	259	6414	5390	6940	-394	-806	-1029
ATOM	3866	N	GLU	B	260	40.012	-24.735	11.957	1.00	45.60	N
ANISOU	3866	N	GLU	B	260	5829	4934	6513	-554	-807	-746

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TABLE 3-continued

ATOM	3867	CA	GLU	B	260	39.382	-24.650	13.282	1.00	45.05	C
ANISOU	3867	CA	GLU	B	260	5726	4914	6476	-621	-608	-543 C
ATOM	3868	C	GLU	B	260	39.635	-23.302	13.922	1.00	43.78	C
ANISOU	3868	C	GLU	B	260	5533	4639	6263	-570	-757	-560 C
ATOM	3869	O	GLU	B	260	38.857	-22.876	14.791	1.00	45.12	O
ANISOU	3869	O	GLU	B	260	5560	5060	6424	-617	-751	-485 O
ATOM	3870	CB	GLU	B	260	39.874	-25.745	14.232	1.00	46.43	C
ANISOU	3870	CB	GLU	B	260	5940	4946	6755	-657	-822	-605 C
ATOM	3871	CG	GLU	B	260	41.393	-25.355	14.338	1.00	60.00	C
ANISOU	3871	CG	GLU	B	260	7704	6586	8506	-572	-796	-610 C
ATOM	3872	CD	GLU	B	260	42.059	-26.909	33.413	1.00	62.93	C
ANISOU	3372	CD	GLU	B	260	8132	6856	8924	-543	-81	-714 C
ATOM	3873	OE1	GLU	B	260	41.756	-25.995	12.178	1.00	58.05	O
ANISOU	3873	OE1	GLU	B	260	7557	6313	8301	-535	-334	-312 O
ATOM	3374	OE2	GLU	B	260	42.960	-27.625	13.923	1.00	66.44	O
ANISOU	3874	OE2	GLU	B	260	8606	7184	9443	-520	-821	-699 O
ATOM	3875	N	TYR	B	261	40.738	-22.523	13.560	1.00	41.38	N
ANISOU	3875	N	TYR	B	261	5247	4550	5927	-478	-71	-571 N
ATOM	3376	CA	TYR	B	261	40.925	-21.251	14.108	1.00	381.45	C
ANISOU	3876	CA	TYR	B	261	4843	4261	5504	-435	-6167	-439 C
ATOM	3877	C	TYR	B	261	39.791	-20.295	13.723	1.00	38.27	C
ANISOU	3877	C	TYR	B	261	4772	4309	5401	-443	-063	-489 C
ATOM	3878	O	TYR	B	261	39.623	-19.259	14.347	1.00	34.93	O
ANISOU	3878	O	TYR	B	261	4313	4010	4944	-427	-630	-422 O
ATOM	3879	CB	TYR	B	261	42.261	-23.666	13.729	1.00	37.23	C
ANISOU	3879	CB	TYR	B	261	4713	4104	4930	-342	-623	-514 C
ATOM	3880	CG	TYR	B	261	43.439	-21.293	14.415	1.00	37.28	C
ANISOU	3880	CG	TYR	B	261	48139	4091	5500	-322	-618	-407 C
ATOM	3881	CD1	TYR	B	261	43.485	-21.428	15.785	1.00	30.90	C
ANISOU	3881	CD1	TYR	B	261	4695	3915	.5410	-3561	-621	-414 C
ATOM	3882	CD2	TYR	B	261	44.541	-21.720	13.666	1.00	39.21	C
ANISOU	3882	CD2	TYR	B	261	5031	4192	5674	-261	-502	-563 C
ATOM	3883	CE1	TYR	B	261	44.597	-21.945	10.443	1.00	37.99	C
ANISOU	3883	CE1	TYR	B	261	4861	3963	5612	-329	-019	-392 C
ATOM	3884	CE2	TYR	B	261	45.6.58	-22.244	14.283	1.00	38.54	C
ANISOU	3884	CE2	TYR	B	261	4970	4015	5659	-232	-604	-547 C
ATOM	3885	CZ	TYR	B	261	45.695	-22.308	15.0.50	1.00	41.31	C
ANISOU	3885	CZ	TYR	B	261	53111.	4328	6055	-265	-612	-461 C
ATOM	3386	OH	TYR	B	261	40.822	-22.949	16.212	1.00	41.52	O
ANISOU	3886	OH	TYR	B	261	5362	4264	16150	-243	-014	-447 O

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TABLE 3-continued

ATOM	3887	N	LYS	B	262	03.293	-20.645	12.711	1.00	39.64	N
ANISOU	3887	N	LYS	B	262	4938	4580	5545	-467	-699	-547 N
ATOM	3888	CA	LYS	B	262	137.905	-19.797	12.295	1.00	40.89	C
ANISOU	3888	CA	LYS	B	262	7040	4804	5628	-471	-702	-550 C
ATOM	3889	C	LYS	B	262	36.305	-1	0.719	13.318	1.00	42.26 C
ANISOU	3889	C	LYS	B	262	5167	5072	6613	-540	-713	-481 C
ATOM	3890	O	LYS	B	262	35.108	-18.701	13.399	1.00	4069	O
ANISOU	3890	O	LYS	B	262	4922	4978	5562	-527	-690	-443 O
ATOM	3891	CB	LYS	B	262	137.3130	-20.246	10.	966	1.00	46.22 C
ANISOU	3891	CB	LYS	B	262	5722	5576	0264	-480	-744	-643 C
ATOM	3892	SG	LYS	B	262	38.201	-19.894	9.752	1.00	47.84	C
ANISOU	3892	CG	LYS	B	262	5965	5800	6412	-394	-722	-735 C
ATOM	3893	CG	LYS	B	262	37.418	-20.379	8.548	1.00	57.68	C
ANISOU	3893	CD	LYS	B	262	7207	7097	7513	-415	-773	-737 C
ATOM	3894	CD	LYS	B	262	38.334	-20.719	7.397	1.00	62.90	C
ANISOU	3894	CE	LYS	B	262	7921	7730	8247	-356	-707	-880 C
ATOM	3895	CE	LYS	B	262	37.704	-21.747	6.505	1.00	75.75	C
ANISOU	3895	NZ	LYS	B	262	9559	9352	9873	-404	-830	-990 N
ATOM	3896	NZ	ASP	B	263	361.641	-20.754	14.119	1.00	41.28	N
ANISOU	3896	N	ASP	B	263	.5051	4862	5772	-619	-738	-462 N
ATOM	3897	N	ASP	B	263	35.741	-20.725	15.287	1.00	43.12	N
ANISOU	3897	CA	ASP	B	263	3238	5.121	6026	-489	-738	-383 C
ATOM	3898	CA	ASP	B	263	36.124	-19.643	16.280	1.00	33.37	C
ANISOU	3398	C	ASP	B	263	4698	4532	5476	-645	-686	-299 C
ATOM	3899	C	ASP	B	263	35.245	-19.002	10.904	1.00	38.09	C
ANISOU	3899	C	ASP	B	263	4533	46102	5337	-663	-672	-244 C
ATOM	3900	CB	ASP	B	263	35.7E1	-22.0138	16.053	1.00	44.25	C
ANISOU	3900	CB	ASP	B	263	5407	5146	62.65	-774	-767	-366 C
ATOM	3901	CG	ASP	B	263	35.232	-22.229	15.181	1.00	43.65	C
ANISOU	3901	CG	ASP	B	263	5978	5655	5357	-83148	10	-822 -450 C
ATOM	3902	OD1	ASP	B	263	94.596	-22.972	14.137	1.30	48.14	O
ANISOU	3902	OD1	ASP	B	263	5832	5672	0738	-831	-848	-510 O
ATOM	3903	OD2	ASP	B	263	-24.36.6	15.442	1.90	50.51	O	
ANISOU	3903	OD2	ASP	B	263	5255	5767	7109	-874	-844	-461 O
ATOM	3904	N	SER	B	264	137.433	-19.433	16.511	1.30	35.53	N
ANISOU	3904	N	SER	B	264	4707	4538	5456	-585	-5.57	-239 N
ATOM	3905	CA	SER	B	264	37.918	-13	425	17.350	1.00	35.02 C
ANISOU	3905	CA	SER	B	264	423	4102	4959	-539	-539	-222 C
ATOM	3906	C	SER	B	264	37.717	-17.949	16.775	1.00	30.07	C
ANISOU	3906	C	SER	B	264	3709	3707	41390	-474	-576	-225 C

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TABLE 3-continued

ATOM	3907	O	SER	B	264	37.401	-16.1	56	17.510	1.00	34.30	O
ANISOU	3907	O	SER	B	264	4088	4174	4772	-465	-545	-169	O
ATOM	3908	CB	SER	B	264	39.11L9	-18.557	17.391	1.00	39.18	C	
ANISOU	3908	CB	SER	B	264	4431	4162	5	53	-493	-589	-216
ATOM	3909	OG	SER	B	264	39.555	-19.599	18.737	1.00	44	77	O
ANISOU	3909	OG	SER	B	264	5535	5163	6307	-544	-609	-174	O
ATOM	3910	N	VAL	B	265	37948	-16901	15.482	1.00	32.99	N	
ANISOU	3910	N	VAL	B	265	39.56	3969	4594	-427	-583	-290	N
ATOM	3911	CA	VAL	B	265	37.701	-15.514	14.809	1.00	32.41	C	
ANISOU	3911	CA	VAL	B	265	3874	3997	4444	-3.64	-5.51	-291	C
ATOM	3912	C	VAL	B	265	36.233	-15.222	15.003	1.00	32.52	C	
ANISOU	3912	C	VAL	B	265	13827	4106	4424	-400	-565	-263	C
ATOM	3913	O	VAL	B	265	35.879	-14.096	15.432	1.00	32.52	O	
ANISOU	3913	O	VAL	B	265	3796	4174	4365	-363	-532	-214	O
ATOM	3914	CB	VAL	B	265	38.065	-15.751	13.331	1.00	34.55	C	
ANISOU	3914	CB	VAL	B	265	4174	4276	4678	-319	-561	-368	C
ATOM	3915	CG1	VAL	B	265	37.602	-14.523	12.533	1.00	38.90	C	
ANISOU	3915	CG1	VAL	B	265	4703	4935	5142	-260	-540	-365	C
ATOM	3916	CG2	VAL	B	265	39.504	-15.383	13.132	1.00	30.28	C	
ANISOU	3916	CG2	VAL	B	265	3685	3645	4164	-269	-532	-388	C
ATOM	3917	N	GLN	B	266	35.350	-16.155	14.752	1.30	32.13	N	
ANISOU	3917	N	GLN	B	266	3882	4182	4523	-470	-615	-295	N
ATOM	3918	CA	GLN	B	266	33.922	-15.958	-5.05	1.00	313.3.9	C	
ANISOU	3918	CA	GLN	B	266	3871	4332	4553	-515	-533	-271	C
ATOM	3919	C	GLN	B	266	39.552	-15.504	16.455	1.00	25.95	C	
ANISOU	3919	C	GLN	B	266	3251	3759	3901	-545	-604	-189	C
ATOM	3920	O	GLN	B	266	32.81313	-14.550	16.597	1.00	30.26	O	
ANISOU	3920	O	GLN	B	266	3368	4016	4113	-523	-583	-154	O
ATOM	3921	CB	GLN	B	266	33.078	-17.176	14.5813	1.00	33.02	C	
ANISOU	3921	CB	GLN	B	266	37119	4250	4517	-6100	-694	-321	C
ATOM	3922	CG	GLN	B	266	611.573	-16.829	14.602	1.00	35.33	C	
ANISOU	3922	CG	GLN	B	266	40513	4722	4840	-536	-713	-306	C
ATOM	3923	CD	GLN	B	266	30.721	-17.964	14.025	1.00	38.13	C	
ANISOU	3923	CD	GLN	B	266	4319	5013	5157	-722	-777	-366	C
ATOM	3924	OE1	GLN	B	266	41136	-13.491	12.938	1.05	43.91	O	
ANISOU	3924	OE1	GLN	B	266	5095	5715	5382	-7113	-810	-442	O
ATOM	3925	NE2	GLN	B	266	29.690	-13.356	14.734	1.00	33.76	N	
ANISOU	3925	NE2	GLN	B	266	4333	5120	5268	-307	-793	-834	N
ATOM	3926	N	GLU	B	267	34.021	-15.392	17.407	1.53	30.37	N	
ANISOU	3926	N	GLU	B	267	3455	3461	4234	-1595	-604	-151	N

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ATOM	3927	CA	GLU	B	267	33.510	-151.182	16.616	1.00	30.19	C
ANISOU	3927	OA	GLU	B	267	3432	31345	4225	-15.33	-580	-32 C
ATOM	3928	C	GLU	B	267	34.203	-14.859	19.356	1.00	30.70	C
ANISOU	3928	C	GLU	B	267	3473	3941	4231	-557	-5215	-40 C
ATOM	3929	O	GLU	B	267	93.537	-14.150	20.163	1.00	32.75	O
ANISOU	3929	O	GLU	B	267	3689	4270	4485	-559	-495	O
ATOM	3930	CB	GLU	B	267	34.1817	-17.333	19.537	1.33	35.10	C
ANISOU	3930	CB	GLU	B	267	4072	48169	4960	-693	-592	-518 C
ATOM	3931	CG	GLU	B	267	1148	10148	1021	313	1.00	37.37 C
ANISOU	3931	CG	GLU	B	267	4315	45.171	5211	-737	-557	27 C
ATOM	3932	CD	GLU	B	267	3.615-485	21.895	1.00	52.51	C	
ANISOU	3932	CD	GLU	B	267	521	-1502	71	4327	-592	711.1 C
ATOM	3933	OE1	GLU	B	267	34.474	-19.388	21.584	1.00	5174	O
ANISOU	3933	OE1	GLU	B	267	6211	62.95	7152	-831	-518	30 O
ATOM	3934	OE2	GLU	B	267	32.786	-18.030	22.835	1.00	73.14	O
ANISOU	3934	OE2	GLU	B	267	161535	5955	7323	-892	-5313	117 O
ATOM	3935	N	THR	B	268	135.42	-14.795	19.042	1.30	26.291	N
ANISOU	3935	N	THR	B	268	8021	3581	3047	-495	-507	-58 N
ATOM	3936	CA	THR	B	268	35.034	-13.2.5	19.511	1.00	29.481	C
ANISOU	3936	CA	THR	B	268	3371	3753	4058	-427	-455	-24 C
ATOM	3937	C	THR	B	268	35.2611	-12.099	13.901	1.00	30.05	C
ANISOU	3937	C	THR	B	268	8470	3993	4134	-375	-435	-27 C
ATOM	3998	O	THR	B	268	136.106	11.061	1	9.555	1.00	2057 O
ANISOU	3998	O	THR	B	268	3460	4045	4111	-344	-396	13 O
ATOM	3939	CB	THR	B	268	37.537-13.117	19.251	1.00	27.08	C	
ANISOU	3939	CB	THR	B	268	31.130	3385	3774	-370	-434	-43 C
ATOM	3940	OG1	THR	B	268	37.869	-13.3135	17.844	1.00	25.24	O
ANISOU	3940	OG1	THR	B	268	2920	3143	3527	-337	-449	-106 O
ATOM	3941	OG2	THR	B	268	138.962	-14.131	20.027	1.00	25.15	O
ANISOU	3941	OG2	THR	B	268	9044	3171	13719	-467	-470	-28 O
ATOM	3942	N	ALA	B	269	34.781	-12.240	17.573	1.00	30.55	N
ANISOU	3942	N	ALA	B	269	34015	3972	4044	-354	-463	-75 N
ATOM	3943	CA	ALA	B	269	33.526	-11.193	17.072	1.00	30.17	C
ANISOU	3943	CA	ALA	B	269	3378	4090	13995	-315	-4.73	-73 C
ATOM	3944	C	ALA	B	269	32.572	-11.072	17.783	1.00	31.29	C
ANISOU	3944	C	ALA	B	269	3441	4302	4122	-655	-458	-37 C
ATOM	3945	O	ALA	B	269	32.022	-9.924	17.945	1.00	32.45	O
ANISOU	3945	O	ALA	B	269	3553	4532	4236	-306	-428	-8 O
ATOM	3946	CB	ALA	B	269	33.730-11476	15.590	1.00	29.56	C	
ANISOU	3946	CB	ALA	B	269	3319	4051	3899	-295	-438	-133 C

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TABLE 3-continued

ATOM	3947	N	GLU	B	270	32.021	-12.161	15.249	1.00	33.60	N
ANISOU	3947	N	GLU	B	270	3717	4085	4465	-445	-492	-36 N
ATOM	3948	CA	GLU	B	270	30.630	-12.152	15.680	1.00	31.261	C
ANISOU	3948	CA	GLU	B	270	3024	4389	4184	-495	-497	-4 C
ATOM	3949	C	GLU	B	270	133.725	-11.567	20.217	1.00	30.90	C
ANISOU	3949	C	GLU	B	270	12.23	8	273	3235	4633	4119 -459 -440 75 C
ATOM	3950	O	GLU	B	270	25.798	-10.6138	20.573	1.00	27.91	O
ANISOU	3950	O	GLU	B	270	2851	4044	3709	-74	-429	32 O
ATOM	3951	CB	GLU	B	270	30.068	-13.518	36.969	1.00	3175	C
ANISOU	3951	CB	GLU	B	270	3352	4404	4276	-603	-545	-19 C
ATOM	3952	CG	GLU	B	270	29.720	-14.1133	17.508	1.05	313	90 C
ANISOU	3952	CG	GLU	B	270	4283	5325	5173	-614	-799	-90 C
ATOM	3953	CD	GLU	B	270	20.241	-15.549	17.5.17	1.00	42.89	C
ANISOU	3953	CD	GLU	B	270	4765	8777	5729	-724	-645	-115 C
ATOM	3954	OE1	GLU	B	270	29.662	-15.421	18.4761	1.00	50.55	O
ANISOU	3954	OE1	GLU	B	270	5774	6668	5763	-731	-644	-86 O
ATOM	3955	OE2	GLU	B	270	26.487	-15.039	15.753	1.00	46.06	O
ANISOU	3955	OE2	GLU	B	270	5217	6307	6203	-754	-694	-155 O
ATOM	3956	N	VAL	B	271	33.751	-11.832	21.010	1.00	28.53	N
ANISOU	3956	N	VAL	B	271	31041	3046	13852	-499	-433	79 N
ATOM	3957	CA	VAL	B	271	21.742	-11.364	22.362	1.30	26.43	C
ANISOU	3957	CA	VAL	B	271	3016	3952	3536	-504	-390	334 C
ATOM	3958	C	VAL	B	271	31.9.35	-9.343	22.540	1.00	28.37	C
ANISOU	3958	C	VAL	B	271	2974	3946	3744	-41 5	-341	147 C
ATOM	3959	O	VAL	B	271	31.882	-9.335	23.671	1.00	32.50	O
ANISOU	3959	O	VAL	B	271	3523	4531	4296	-4 4	-305	186 O
ATOM	3960	CB	VAL	B	271	32.721	-12	143	23.267	1.00	28.17 C
ANISOU	3960	CB	VAL	B	271	30134	3820	13849	-545	-391	150 C
ATOM	3961	CG1	VAL	B	271	32.373	-13.613	23.217	1.00	31.50	C
ANISOU	3961	CG1	VAL	B	271	3455	4208	41333	-636	-436	156 C
ATOM	3962	CG2	VAL	B	271	34.174	-11.906	22.773	1.00	20.93	C
ANISOU	3962	CG2	VAL	B	271	2945	3580	3703	-484	-385	132 C
ATOM	3963	N	VAL	B	272	32.451	-9.147	21.524	1.00	26.36	N
ANISOU	3963	N	VAL	B	272	2746	3689	3463	-343	-335	115 N
ATOM	3964	CA	VAL	B	272	32079	-7.711	21.644	1.00	29.19	C
ANISOU	3964	CA	VAL	B	272	3145	4120	3824	-26.3	-288	129 C
ATOM	3965	C	VAL	B	272	31.355	-6.952	21.055	1.00	29.92	C
ANISOU	3965	C	VAL	B	272	3181	4315	3873	-221	-287	127 C
ATOM	3966	O	VAL	B	272	31	285	-5.707	2	104	1.00 30.40 O
ANISOU	3966	O	VAL	B	272	3243	44001	0902	-149	-248	140 O

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TABLE 3-continued

ATOM	3967	CB	VAL	B	272	33.896	-7.224	20.992	1.00	29.75	C
ANISOU	3967	CB	VAL	B	272	3282	412.5	389.5	-204	-271	107 C
ATOM	3968	CG1	VAL	B	272	35.342	-7.847	21.739	1.30	31.37	C
ANISOU	3968	CG1	VAL	B	272	3532	4243	4144	-239	-270	115 C
ATOM	3969	CG2	VAL	B	272	33.903	-7.519	19.527	1.00	32.01	C
ANISOU	3969	CG2	VAL	B	272	3555	4497	4245	-188	-305	54 C
ATOM	3970	N	ILE	B	273	30.365	-7.558	20.542	1.00	29.31	N
ANISOU	3970	N	ILE	B	273	3053	4288	3795	-288	-330	112 N
ATOM	3971	CA	ILE	B	273	29.055	-7.074	20.106	1.00	29.53	C
ANISOU	3971	CA	ILE	B	273	3010	4428	13783	-232	-335	114 C
ATOM	3972	C	ILE	B	273	28.437	-6.214	21.234	1.00	30.42	C
ANISOU	3972	C	ILE	B	273	3459	4977	4231	-211	-288	154 C
ATOM	3973	O	ILE	B	273	27.987	-5.076	23.977	1.00	35.43	O
ANISOU	3973	O	ILE	B	273	3599	5297	4480	-131	-255	181 O
ATOM	3974	CB	ILE	B	273	28.032	-8.076	19.610	1.30	32.26	C
ANISOU	3974	CB	ILE	B	273	3294	4827	4137	-301	-350	92 C
ATOM	3975	CG1	ILE	B	273	28.457	-3.719	18.300	1.30	33.46	C
ANISOU	3975	CG1	ILE	B	273	3484	4941	4288	-333	-437	40 C
ATOM	3976	CG2	ILE	B	273	28.679	-7.355	19.255	1.00	38.02	C
ANISOU	3976	CG2	ILE	B	273	3941	5679	4824	-259	-394	97 C
ATOM	3977	CD1	ILE	B	273	26.30	-7.743	17.227	1.00	85.44	C
ANISOU	3977	CD1	ILE	B	273	17771	5233	4490	-203	-4261	25 C
ATOM	3978	N	PRO	B	274	28.428	-6.729	22.470	1.00	33.29	N
ANISOU	3978	N	PRO	B	274	3433	4944	4270	-276	-274	19 N
ATOM	3979	CA	PRO	B	274	27.844	-5.927	23.513	1311	35.89	C
ANISOU	3979	CA	PRO	B	274	3723	5335	4580	-253	-228	213 C
ATOM	3980	C	PRO	B	274	28.644	-4.634	23.779	1.00	38.47	C
ANISOU	3980	C	PRO	B	274	4101	5828	4886	-165	-177	217 C
ATOM	3981	O	PRO	B	274	28.210	-3.774	24.306	1.00	34.70	O
ANISOU	3981	O	PRO	B	274	3598	5197	4388	-131	-135	236 O
ATOM	3982	CB	PRO	B	274	27.934	-8.844	24.716	1.00	34.49	C
ANISOU	3982	CB	PRO	B	274	3544	5131	4431	-344	-223	242 C
ATOM	3983	CG	PRO	B	274	27.878	-3.234	24.152	1.00	36.37	C
ANISOU	3983	CG	PRO	B	274	3750	5332	4700	-423	-278	227 C
ATOM	3984	CD	PRO	B	274	28.8013	-8.071	22.997	1.00	35.74	C
ANISOU	3984	CD	PRO	B	274	3763	5191	4827	-373	-293	1821 C
ATOM	3985	N	LEU	B	275	29.810	-4.452	23.148	1.30	32.71	N
ANISOU	3985	N	LEU	B	275	2.444	48121	4165	-133	-180	199 N
ATOM	3986	CA	LEU	B	275	30.076	-3.256	23.372	1.00	35.50	C
ANISOU	3986	CA	LEU	B	275	3871	5149	4528	-61	-134	230 C

TABLE 3-continued

ATOM	3987	C	LEU	B	275	30.477	-2.184	22.280	1.00	35.47	13	C	
ANISOU	3987	C	LEU	B	275	3973	5278	4531	131	-123	131	C	
ATOM	3988	O	LEU	B	275	31.124	-1.099	22.1315	1.00	35.49	0		
ANISOU	3988	O	LEU	B	275	3899	5115	44709.5	-83	1148	103	O	
ATOM	3989	CB	LEU	B	275	32.183	-3.645	23.502	1.00	29.25	C		
ANISOU	3989	CB	LEU	B	275	3145	4245	3763	-82	-135	191	C	
ATOM	3990	OG	LEU	B	275	32.506	-4.354	24.411	1.00	34.71	0		
ANISOU	3990	OG	LEU	B	275	.3821	4887	4471	-1	39	-155	204	O
ATOM	3991	CD1	LEU	B	275	34.005	-5.142	24.536	1.06	28.38	C		
ANISOU	3991	CD1	LEU	B	275	3095	3988	3730	-177	-1.57	19.5	C	
ATOM	3992	CD2	LEU	B	275	31.858	-4.586	25.772	1.00	28.73	C		
ANISOU	3993	CD2	MET	B	276	3034	4181	3732	-195	-129	234	C	
ATOM	3993	N	MET	B	276	29.571	-2.489	21.353	1.00	35.93	N		
ANISOU	3993	N	MET	B	276	3834	5273	4515	30	-150	SC	N	
ATOM	3994	CA	MET	B	276	29.158	-1.591	23.28.5	1.30	37.45	C		
ANISOU	3994	CA	MET	B	276	4051	5507	4671	121	-159	181	C	
ATOM	3995	C	MET	B	276	28.081	-0.345	20.735	1.00	43	91	C	
ANISOU	3995	C	MET	B	276	4432	6027	5.336	177	-132	202	C	
ATOM	3996	O	MET	B	276	27.291	-0.962	21.638	1.00	35.36	O		
ANISOU	3996	O	MET	B	276	3659	6.378	4390	135	-128	21	O	
ATOM	3997	CB	MET	B	276	28.587	-2.413	191387	1.00	34.08	C		
ANISOU	3997	CB	MET	B	276	3574	5195	4306	99	-219	158	C	
ATOM	3998	CG	MET	B	276	29.801	-3.220	18.450	1.00	34.54	C		
ANISOU	3998	CG	MET	B	276	3720	5097	4357	65	-242	130	C	
ATOM	3999	SD	MET	B	276	31.405	-2.358	18.159	1.0036.59	S			
ANISOU	3999	SD	MET	B	276	4127	5308	4519	126	-195	532	S	
ATOM	4000	CE	MET	B	276	32.513	-19754	18.405	1.00	44.55	C		
ANISOU	4000	CE	MET	B	276	5127	6176	5325	40	-218	1135	C	
ATOM	4001	N	ALA	B	277	2-.988	0.698	20.094	1.00	38.41	N		
ANISOU	4001	N	ALA	B	277	4139	5729	4750	275	-111	211	N	
ATOM	4002	CA	ALA	B	277	26.927	1.549	20.535	1.50	39.22	C		
ANISOU	4002	CA	ALA	B	277	4179	5899	41025	344	-82	230	C	
ATOM	4003	C	ALA	B	277	25.479	0.994	20.372	1.00	40.89	C		
ANISOU	4003	C	ALA	B	277	4286	6229	5022	321	-121	230	C	
ATOM	4004	O	ALA	B	277	25.326	3.196	19.422	1.00	38.58	O		
ANISOU	4004	O	ALA	B	277	3960	5958	4722	288	-175	215	O	
ATOM	4005	CB	ALA	B	277	27.073	2.189	19.545	1.00	34.94	C		
ANISOU	4005	CB	ALA	B	277	3.575	5343	4258	4.53	-61	243	C	
TEF	4306	ALA	B	277									
HETATM	4007	C1	BMX	A1000		35.951	-4.950	18.275	0.40	12.00	C		

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TABLE 3-continued

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HETATM 4008 C2 BMX A1000	36.393 -5.915 17.210 0.40 12.88 C
HETATM 4009 C3 BMX A1000	37.834 -5.506 16.1356 0.40 12.35 C
HETATM 4010 C4 BMX A1000	37.896 -4.121 16.436 0.40 12.97 C
HETATM 4011 C5 BMX A1000	37.430 -3.163 17.578 0.40 12.35 C
HETATM 4012 C6 BMX A1000	37.3E2 -1.356 17.242 0.40 1 1.98 C
HETATM 4013 C7 BMX A1000	30.212 -0.507 15.280 0.40 13.34 C
HETATM 4014 C8 BMX A1000	34.190 -5.327 14.150 0.40 15.65 C
HETATM 4015 N2 BMX A1000	35.427 -5.693 151078 3.40 12.59 N
HETATM 4016 O1 BMX A1000	35.677 -5.278 19.503 0.40 12.10 O
HETATM 4017 O3 BMX A1000	38.276 -3.497 15.843 3.43 12.50 O
HETATM 4018 O4 BMX A1000	39.211 -3.784 16.0313 0.40 12.01 O
HETATM 4019 O5 BMX A1000	96.124 -3.574 17.984 0.40 13.50 O
HETATM 4020 O6 BMX A1000	.35.576 -1.4841 15.032 0.40 12.52 O
HETATM 4021 O7 BMX A1000	35.7.82 -7.664 15.426 0.40 14.76 O
HETATM 4022 P BMX A1000	36.529 -3.381 15.428 0.40 12.70 P
HETATM 4023 O19 BMX A1000	35.473 -0.222 14.347 0.40 12.53 O
HETATM 4024 O17 BMX A1000	36.329 0.13.1 8 16.500 0.40 13.63 O
HETATM 4025 O18 BMX A1000	37.856 0.035 14.833 0.40 13.15 O
HETTAM 4026 C1 BMX B1010	64.385 3.763 17.8.49 0.40 13.55 C
HETATM 4027 C2 BMX B1000	63.911 4.6213 16.704 0.40 113.45 C
HETATM 4028 C3 BMX B1000	132.489 4.221 10.321 0.40 12.62 C
HETATM 4029 C4 BMX B1000	62.371 2.849 151073 0.40 12.57 C
HETATM 4030 C5 BMX B1000	62. 780 2.025 17.191 0.40 11.53 C
HETATM 4031 C6 BMX B1000	52.780 0.467 17.35 0.40 12.40 C
HETATM 4032 C7 BMX B1000	05.096 5.076 14.609 0.40 14.20 C
HETATM 4033 C8 BMX B1000	66.104 4.645 133.590 0.40 14. 11 C
HETATM 4034 N2 BMX B1000	64.951 4.326 15.60.6 0.40 13.41 N
HETATM 4035 O1 8MX 81000	53.605 4.171 39.029 0.12.52 O
HETATM 4036 O3 BMX B1000	82. I1 515 4.968 15.140 0.40 11.00 O
HETATM 4037 O4 BMX B1000	61.090 2002 15.532 040 12.142 O
HETATM 4038 O5 BMX B1000	94. I 39 2.397 17.4619 0.40 14.10 O
HETATM 4039 O6 BMX B1000	63.755 0.045 16.220 0.40 14.36 O
HETATM 4040 O7 BMX B1000	64.440 6.104 14.426 0.46 14.53 O
HETATM 4041 P BMX B1000	53.612 -1.431 15.592 0.40 13.06 P
HETATM 4042 O19 BMX 81000	54.055 -2.214 16.804 0.40 12.42 O
HETATM 4043 O17 BMX B1000	02.520 -1.596 14.878 0.40 13.41 O
HETATM 4044 O18 BMX B1000	614.846 -1367 14.515 0.40 14.95 O
HETATM 4045 O HOH A1101	30.465 -0.098 29.486 1.00 28.80 O
HETATM 4046 O HOH A1102	39.030 -2.218 13.942 1.00 05.02 O
HETATM 4047 O HOH A1103	38.435 1.950 18. 1 39 1.00 27.90 O

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TABLE 3-continued

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HETATM 4048 O HOH A1104	41.206	15.194	32.047	1.00	29.52	O
HETATM 4049 O HOH A1105	24.122	-3187	45.575	1.00	32.50	O
HETATM 4050 O HOH A1106	28.552	5.104	4.7813	1.00	44.36	O
HETATM 4051 O HOH A1107	39.011	0.578	28.874	1.00	30.12	O
HETATM 4052 O HOH A1108	34.971	-2.503	12.953	1.00	133.62	O
HETATM 4053 O HOH A1109	35.717	-9.56.5	13.356	1.00	31.00	O
HETATM 4054 O HOH A1110	41.621	-0.576	12.205	1.00	30.37	O
HETATM 4055 O HOH A1111	31.410	-0.020	0.502	1.00	35.17	O
HETATM 4059 O HOH A1112	58.166	8.145	8.5.38	1.00	34.35	O
HETATM 4057 O HOH AA113	56.635	7.216	6.355	1.00	30.85	O
HETATM 4058 O HOH AA114	67.208	-0.418	23.775	1.00	30.20	O
HETATM 4059 O HOH A1115	23.474	-1.938	13.231	1.00	39.56	O
HETATM 4060 O HOH A4116	41.617	1.441	12.532	1.00	29.73	O
HETATM 4061 O HOH A1117	43.151	-8.191	4.13	1.00	37.99	O
HETATM 4062 O HOH A1118	34.754	6.476	27.195	1.00	35.64	O
HETATM 4063 O HOH A1119	66.715	7.574	3.049	1.00	41.41	O
HETATM 4064 O HOH A1120	30.255	1.006	28.320	1.00	33.01	O
HETATM 4065 O HOH A1121	32.403	-8.000	12.153	1.00	34.17	O
HETATM 4066 O HOH A1122	57.669	20.047	16.073	1.00	36.58	O
HETATM 4067 O HOH A1123	56.846	14.757	21.072	1.00	32.45	O
HETATM 4068 O HOH A1124	53.117	4.579	19.863	1.00	32.03	O
HETATM 4069 O HOH A1125	52.710	19.584	20.978	1.00	33.28	O
HETATM 4070 O HOH A1126	45.251	4.092	-6.444	1.00	45.84	O
HETATM 4071 O HOH A1127	24.099	0.043	8719	1.00	32.03	O
HETATM 4072 O HOH A1128	31.887	17.106	27.223	1.00	37.16	O
HETATM 4073 O HOH A1129	27.750	-5.905	34.060	1.00	34.00	O
HETATM 4074 O HOH A1130	41.985	23.528	20.633	1.00	37.00	O
HETATM 4075 O HOH A1131	35.617	2.935	2071.00	40.86	O	
HETATM 4076 O HOH 01132	43.640	22.266	16266	1.00	46.36	O
HETATM 4077 O HOH A1133	59.071	14.463	20.497	1.30	32.43	O
HETATM 4078 O HOH A1134	23.153	-4.551	19.068	1.00	39.58	O
HETATM 4079 O HOH A1135	35.5231	-6.086	5.61731	1.00	36.75	O
HETATM 4080 O HOH A1136	310.142	-0.007	0.105	1.00	43.31	O
HETATM 4081 O HOH A1137	43.2134	3.553	10.847	1.00	32.43	O
HETATM 4082 O HOH A1138	74.313	12.8137	1499.5	1.00	51.72	O
HETATM 4083 O HOH A1139	55.508	11.245	11.765	1.00	33.81	O
HETATM 4084 O HOH A1140	30.370	-1.050	11.032	1.00	38.06	O
HETATM 4085 O HOH A1141	42.445	2.13.92	9.635	1.00	44.35	O
HETATM 4086 O HOH A1142	38.461	110.587	3.2141	1.00	40.74	O
HETATM 4087 O HOH A1143	20.9115	13.4154	1.00	34.21	O	

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TABLE 3-continued

HETATM 4088 O HOH A1144	53.546 21.573 19.214 1.06 44.07 O
HETATM 4089 O HOH A1145	1134.057 20.084 1 84502 1.190 38.13 O
HETATM 4090 O HOH A1146	34.928 20.835 6.690 1.00 43.83 O
HETATM 4091 O HOH A1147	47.275 22.301 5.1 23 1.00 38.67 O
HETATM 4092 O HOH A1148	38.435 -1.1.335 11 .2915 1.00 40.65 O
HETATM 4093 O HOH A1149	65.101 14.753 9.176 1.00 44.42 O
HETATM 4094 O HOH A4150	55.167 5.379 2.528 LOC 48.95 O
HETATM 4095 O HOH A1151	Si 967 118.590 5.654 1.00 58.83
HETATM 4096 O HOH A1152	58.137 13.747 8.963 1.00 38.1 O
HETATM 4097 O HOH A1153	57.028 14.677 3.327 101 44.19 O
HETATM 4098 O HOH A1154	39.073 25.399 26.3.55 1.00 51.90 O
HETATM 4099 O HOH 41155	21.205 -0.650 17.748 1.00 37.56 O
HETATM 4100 O HOH A1156	53.905 22.451 15.518 1.00 46.38 O
HETATM 4101 O HOH A1157	60.311 20.554 14.377 1.00 54.87 O
HETATM 4102 O HOH A1158	45 175 22.841 25.078 1.00 48.02 O
HETATM 4103 O HOH A1159	35.1351 -13.551 8.451 1.00 50.40 O
HETATM 4104 O HOH A1160	30.036 -5.172 4.501 1.00 34.43 O
HETATM 4105 O HOH A1161	28.780 5.316 28.446 1.00 40.01 O
HETATM 4106 O HOH A1162	5.05E 11002 5.327 1.00 47.15 O
HETATM 4107 O HOH A1163	13.551 1.784 18.231 1.00 53.17 O
HETATM 4108 O HOH A1164	36.586 16.314 E.E22. 1.00 44.73 O
HETATM 4109 O HOH A1165	58 632 24.913 8.912 1.00 51.32 O
HETATM 4110 O HOH A1166	32.254 -9.40.6 4.077 1.00 43.27 O
HETATM 4111 O HOH A1167	13.130 -1.308 0.926 1.00 52.06 O
HETATM 4112 O HOH A1168	39.255 -10.485 6.207 1.00 41.34 O
HETATM 4113 O HOH A1169	42.040 0.213 3.713 1.00 42.51 O
HETATM 4114 O HOH A1170	20.276 4.602 17.070 1.00 52.01 O
HETATM 4115 O HOH A1171	24.227 -7.525 -3.4 79 1.00 58.07 O
HETATM 4116 O HOH A1172	68 290 6.836 14.687 1.00 36.25 O
HETATM 4117 O HOH A1173	28.488 -3.131 14.77.5 1.00 47.26 O
HETATM 4118 O HOH A1174	77.346 3.633 18.650 1.00 47.51 O
HETATM 4119 O HOH A1175	54.010 12.709 23.603 1.00 42.75 O
HETATM 4120 O HOH A1176	25.791 -7.495 -3.071 1.00 62.43 O
HETATM 4121 O HOH A1177	26.392 13.046 17.978 1.00 50.24 O
HETATM 4122 O HOH A1178	40.114 15.398 -2.954 1.00 52.69 O
HETATM 4123 O HOH A1179	22.281 2.038 17.799 1.00 57.68 O
HETATM 4124 O HOH A1180	39.580 -10.265 0.165 1.00 40.17 O
HETATM 4125 O HOH 41181	78.174 -2.251 22.270 1.00 63.98 O
HETATM 4126 O HOH 41182	55.579 21.061 17.087 1.00 54.36 O
HETATM 4127 O HOH A1188	49.560 2.32.6 -6.677 1.50 54.33 O

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TABLE 3-continued

HETATM 4128 O HOH A1134	10.605	6.963	6.381	1.00	58.45	O
HETATM 4129 O HOH A1185	27.486	-0.031	1.856	1.00	58.54	O
HETATM 4130 O HOH A1186	17.432	4.022	18.027	1.00	53.25	O
HETATM 4131 O HOH A1187	28.145	-1.454	12.645	1.00	47.59	O
HETATM 4132 O HOH A1388	15.826	3.344	155.3	1.00	54.47	O
HETATM 4133 O HOH A1189	76.534	5.024	29.679	1.00	52.40	O
HETATM 4134 O HOH A1190	29.670	-0.501	3.280	1.00	54.57	O
HETATM 4135 O HOH A1191	34.828	-4.100	2.643	1.00	54.50	O
HETATM 4136 O HOH A1192	17.875	8.877	11.913	1.00	57.82	O
HETATM 4137 O HOH A1193	30.703	4.390	13.614	1.00	50.75	O
HETATM 4138 O HOH A1194	150.71	-12.775	21.159	1.00	56.73	O
HETATM 4139 O HOH A1195	15.747	-4.073	23.775	1.00	57.54	O
HETATM 4140 O HOH A1196	23.5.52	6.396	10.25.5	1.00	50.77	O
HETATM 4141 O HOH A1197	72.578	14.721	11.504	1.00	53.18	O
HETATM 4142 O HOH A1198	50.930	-10.094	-4.530	1.08	61.96	O
HETATM 4143 O HOH A1199	31.293	-3.475	14.65.8	3 .00	43.66	O
HETATM 4144 O HOH A1200	74.7	1123.574	20.512	1.00	52.16	O
HETATM 4145 O HCH A1201	28.322	8.300	24.68	1.00	53.68	O
HETATM 4146 O HOH A1202	6.272	14228	1 2.21	71.30	60.57	O
HETATM 4147 O HOH A1203	15.763	7.723	15.403	1.00	53.24	O
HETATM 4148 O HOH A1204	45.842	20.652	27.74.5	1.00	48.24	O
HETATM 4149 O HOH A1205	1034	7.319	17 .647	1.00	49.01	O
HETATM 4150 O HOH A1206	70.204	-0.047	133.040	1.00	53.02	O
HETATM 4151 O HOH A1207	26.170	10.770	8.655	1.00	53.05	O
HETATM 4152 O HOH A1208	29.139	-5.559	-0.384	1.00	57.78	O
HETATM 4153 O HOH A1209	31.24-7	3.811	-4073	1.00	58.20	O
HETATM 4154 O HOH A1210	1.208E-i	8.783	35.359	1.60	51.55	O
HETATM 4155 O HOH A1211	73.371	16.646	15.914	1.00	63.09	O
HETATM 4156 O HOH A1212	37.392	2000	14.789	1.00	52.80	O
HETATM 4157 O HOH A1213	132.554	-3.592	1.350	1.00	57.00	O
HETATM 4158 O HOH B1101	43.445	-12.933	13.130	1.00	30.77	O
HETATM 4159 O HOH B1102	61.957	-3.228	18.426	1.00	27.72	O
HETATM 4160 O HOH B1103	56.644	7.580	11.066	1.00	2055	O
HETATM 4161 O HOH B1104	64.584	7.679	12.162	1.00	26.43	O
HETATM 4162 O HOH B1105	68.294	7.048	7.1 18	1.00	39.24	O
HETATM 4163 O HOH B1106	38.148	0.065	23.727	1.00	28.31	O
HETATM 4164 O HOH B1107	88.593	0.080	-4.686	1.00	40.19	O
HETATM 4165 O HOH B1108	54.350	-20.633	30.086	1.00	42.25	O
HETATM 4166 O HOH B1109	58024	-16.951	33.625	1.00	38.70	O
HETATM 4167 O HOH B1110	48.893	-9.716	7.282	1.00	31.70	O

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TABLE 3-continued

HETATM 4168 O HOH B1111	73.605 -2.672 11 3.724 1.00 35.48 O
HETATM 4169 O HOH B1112	47.346 -5.509 20.446 1.00 30.24 O
HETATM 4170 O HOH B1113	65.410 01710 12.765 1.00 29.59 O
HETATM 4171 O HOH B1114	42.218 -15.442 8.515 1.00 40.01 O
HETATM 4172 O HOH B1115	59.595 10.150 11.257 1.00 31.84 O
HETATM 4173 O HOH B1116	61.405 6.525 13.733 1.00 29.38 O
HETATM 4174 O HOH B1117	61.288 -0.564 28.141 1.00 32.31 O
HETATM 4175 O HOH B1118	73.783 10284 4 508 1.50 29.90 O
HETATM 4176 O HOH B1119	66.299 1.756 15.268 1.00 30.10 O
HETATM 4177 O HOH B1120	68.906 -1329 3.525 1.00 39.52 O
HETATM 4178 O HOH B1121	58.023 -3.308 12.958 1.00 35.26 O O
HETATM 4179 O HOH B1122	63.854 -0.357 25.535 1.00 27.81 O
HETATM 4180 O HOH B1123	69.335 1.910 14.691 1.00 37.18 O
HETATM 4181 O HOH B1124	39.732 -22.286 17.526 1.00 51.75 O
HETATM 4182 O HOH B1125	66.367 -2.401 8.678 1.00 31.75 O
HETATM 4183 O HOH B1126	72.632 4.309 13.519 1.50 36.19 O
HETATM 4184 O HOH B1127	63.846 -18.881 9.336 1.00 44.52 O
HETATM 4185 O HOH B1128	45.644 -0.269 3.608 1.00 42.45 O
HETATM 4186 O HOH B1129	71.416 -8.351 5.550 1.00 44.26 O
HETATM 4187 O HOH B1130	40.923 -12.250 12.628 1.00 30.58 O
HETATM 4188 O HOH B1131	69.872 4.518 13.787 1.00 34.81 O
HETATM 4189 O HOH B1132	50.304 -5.522 25.727 1.00 59.39 O
HETATM 4190 O HOH B1133	69.135 -24.163 25.129 1.00 36.44 O
HETATM 4191 O HOH B1134	65.221 -21.288 23.182 1.00 39.91 O
HETATM 4192 O HOH B1135	42.215 -13.132 10.462 1.00 39.94
HETATM 4193 O HOH B1136	86.594 -2.855 16712 1.00 56.19 O
HETATM 4194 O HOH B1137	43.508 -15.265 23.787 1.00 33.71 O
HETATM 4195 O HOH B1138	46.883 -22.512 21.907 1.00 46.45 O
HETATM 4196 O HOH B1139	71.612 -6.230 23.991 1.00 40.05 O
HETATM 4197 O HOH B1140	85.465 -1.039 -1 331 1.00 50.57 O
HETATM 4198 O HOH B1141	64.878 3.401 4.887 1.00 41.08 O
HETATM 4199 O HOH 91142	72.251 -9.109 25.995 1.00 48.37 O
HETATM 4200 O HOH B1143	61.795 9.075 10.217 1.00 41.33 O
HETATM 4201 O HOH B1144	47.5 -20.177 23.304 1.00 35.78 O
HETATM 4202 O HOH B1145	72.25 -5.453 26.533 1.00 47.83 O
HETATM 4203 O HOH B1146	84.575 -21.473 3.479 1.00 48.79 O
HETATM 4204 O HOH B1147	509.95 -26.151 09.529 1.00 54.01 O
HETATM 4205 O HOH B1148	42.278 -17.325 25.240 1.00 37.14 O
HETATM 4206 O HOH B1149	64.141 11.1195 6.974 1.40 41.72 O
HETATM 4207 O HOH B1150	32.737 -19.072 17.314 1.00 44.42 O

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TABLE 3-continued

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HETATM 4208 O HOH B1151	73.40 -12. 59 26.324 1.00 45.95 O
HETATM 4209 O HOH B1152	64.118 -3.623 0.383 1.011 48.28 O
HETATM 4210 O HOH B1153	64.332 -5.449 11.375 1.00 34.16 O
HETATM 4211 O HOH B1154	65.586 -5.1858 28.028 1.00 33.16 O
HETATM 4212 O HOH B1155	25.136 -7.023 11.315 1.00 45.59 O
HETATM 4213 O HOH B1156	65.065 4.530 2.331 1.00 56.64 O
HETATM 4214 O HOH B1157	55.473 0.531 19.553 1.00 37.85 O
HETATM 4215 O HOH B1158	55.433 -9.285 -5.505 1.00 45.37 O
HETATM 4216 O HOH B1159	64.8515 -8.140 3090 1.03 42.78 O
HETATM 4217 O HOH B1160	42.493 -237 19.229 1.00 39.45 O
HETATM 4218 O HOH B1161	40.534-17.709 5.155 1.00 48.87 O
HETATM 4219 O HOH B1162	52.315 -10.548 14.290 1.00 61.77 O
HETATM 4220 O HOH B1163	69.579 -1.052 11.195 1.00 39.94 O
HETATM 4221 O HOH B1164	65058 -25.022 27.155 1.00 58.23 O
HETATM 4222 O HOH B1165	27.364 -20.364 13.978 1.00 58.12 O
HETATM 4223 O HOH B1166	56.743 -3.350 -7.001 1.00 55.60 O
HETATM 4224 O HOH B1167	57.455 5.089 3.767 1.00 42.61 O
HETATM 4225 O HOH B1168	74.410 -11158 14.258 1.00 45.27 O
HETATM 4226 O HOH B1169	75.149 3.405 -1.880 1.00 47.09 O
HETATM 4227 O HOH B1170	73.933 -11.056 15.525 1.00 49.39 O
HETATM 4228 O HOH B1171	60.095 -19.355 -1.096 1.00 60.45 O
HETATM 4229 O HOH B1172	76.550 10.871 15.710 1.00 50.97 O
HETATM 4230 O HOH B1173	65.515 1.051 2.667 1.00 44.41 O
HETATM 4231 O HOH B1174	48.765 -3.773 27.310 1.00 52.46 O
HETATM 4232 O HOH B1175	55.035 1.172 11.890 1.03 35.08 O
HETATM 4233 O HOH B1175	55.580 -2.225 30.146 1.00 38.49 O
HETATM 4234 O HOH B1177	60.982 -22.825 20.125 1.00 60.83 O
HETATM 4235 O HOH B1178	70.874 4.773 1.547 1.00 59.50 O
HETATM 4236 O HOH B1179	33.353 -2.624 2.943 1.00 60.79 O
HETATM 4237 O HOH B1180	45.955 -24.132 18.709 1.00 56.26 O
HETATM 4238 O HOH B1181	74.945 1 .752 -0.840 1.00 56.16 O
HETATM 4239 O HOH B1182	74.922 -13.8003 20.013 1.00 46.41 O
HETATM 4240 O HOH B1183	67.533 0.340 1.045 1.00 54.61 O
HETATM 4241 O HOH B1184	62.224 -24.765 21.501 1.00 53.17 O
HETATM 4242 O HOH B1185	72.114 -0.599 12.716 1.00 45.35 O
HETATM 4243 O HOH B1185	58.409 -17.060 29.630 1.00 36.20 O
HETATM 4244 O HOH B1187	58.269 -24.629 23.224 1.00 46.73 O
HETATM 4245 O HOH B1188	25110 -14575 16.100 1.00 49.70 O
HETATM 4246 O HOH B1189	43.338 -25.893 16.745 1.00 70.93 O
HETATM 4247 O HOH B1190	69.241 -7.602 4.510 1.00 54.36 O

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TABLE 3-continued

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HETATM 4248 O HOH B1191	90.426 -4.832 10.952 1.00 57.23 O
HETATM 4249 O HOH B1192	60.6.40 -1.837 -1.553 1.00 55.15 O
HETATM 4250 O HOH B1193	30.525 -18.750 13.084 1.00 45.99 O
HETATM 4251 O HOH B1194	65.805 -23.265 13.381 1.00 53.74 O
HETATM 4252 O HOH B1195	61.269 -12.59.5 4.931 1.00 54.63 O
HETATM 4253 O HOH B1196	74.321 -5.E.05 2.777 1.00 49.84 O
HETATM 4254 O HOH B1197	76.1.33 15.725 1.187 1.00 60.42 O
HETATM 4255 O HOH B1198	56.275 -23.442 14.182 1.00 43.55 O
HETATM 4256 O HOH B1199	55.157 -24.527 21.55.7 1.00 56.56 O
HETATM 4257 O HOH B1200	73.165 -0.542 23.441 1.00 40.52 O
HETATM 4258 O HOH B1201	58.271 -4.513 3.739 1.00 45.72 O
HETATM 4259 O HOH B1202	98.042 -0.400 0.918 1.00 54.85 O
HETATM 4260 O HDH B1203	76.096 -11.404 4.579 1.00 58.79 O
HETATM 4261 O HOH B1204	70.970 -14.429 11.398 1.00 50.39 O
HETATM 4262 O HOH B1205	97.205 -17.263 31.503 1.00 48.05 O
HETATM 4263 O HOH B1206	58.029 -75.055 5.295 1.00 50.57 O
HETATM 4264 O HOH B1207	53.174 -25.139 8.451 1.00 50.23 O
HETATM 4265 O HOH B1208	77.029 14.1343 13.644 1.00 48.90 O
CONECT 4007 4008 4016 4019	
CONECT 4008 4007 4009 4015	
CONECT 4009 4008 4010 4017	
CONECT 4010 4009 4011 4018	
CONECT 4011 4010 4012 4019	
CONECT 4012 4011 4020	
CONECT 4013 4014 4015 4021	
CONECT 4014 4013	
CONECT 4015 4008 4013	
CONECT 4016 4007	
CONECT 4017 4009	
CONECT 4018 4010	
CONECT 4019 4007 4011	
CONECT 4020 4012 4022	
CONECT 4021 4013	
CONECT 4022 4020 4023 4024 4025	
CONECT 4023 4022	
CONECT 4024 4022	
CONECT 4025 4022	
CONECT 4026 4027 4035 4038	
CONECT 4027 4025 4028 4034	
CONECT 4028 4027 4029 4036	

TABLE 3-continued

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CONECT 4029 4028 4030 4037
CONECT 4030 4029 4031 4038
CONECT 4031 4030 4039
CONECT 4032 4033 4034 4040
CONECT 4033 4032
CONECT 4034 4077 4032
CONECT 4035 4025
CONECT 4036 4028
CONECT 4037 4029
CONECT 4038 4025 4030
CONECT 4039 4031 4041
CONECT 4040 4032
CONECT 4041 4039 4042 4043 4044
CONECT 4042 4041
CONECT 4043 4041
CONECT 4044 4041

MASTER 478 0 2 28 10 0 10 6 4263 2 38 44
END

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**Example 9****Electron Microscopy and Single Particle Analysis**

Purified NanR protein, the NanR/ligand complex, and the NanR/DNA complex were diluted to a final concentration of 300 nM with 300 mM NaCl and 50 mM Tris-HCl (pH 7.0). 5  $\mu$ l of each sample solution were applied to a carbon-coated grid that had been glow-discharged for 3 minutes in air, and the grid was immediately negatively stained using 1% uranyl acetate. Grids were examined in a Technai G2 Spirit Twin transmission electron microscope (FEI, Hillsboro, Oreg., USA), and images were recorded on a 4Kx4K Ultrascan 895 CCD (Gatan, Pleasanton, Calif., USA) at a magnification of 0.36 nm/pixel. For single particle analysis, images of individual particles were selected interactively, windowed out, and imported into the SPIDER program (Health Research, Rensselaer, N.Y., USA). A total of 894 NanR particles, 1131 NanR/ligand particles, and 1039 NanR/DNA particles were used in the processing and class averages were produced by the known reference free method. To compare similar views of the three samples, the datasets were combined and then co-aligned and classified. Averages in which the NanR region was similarly observed were selected for further analysis. The UCSF Chimera program was used for visualization and comparative analyses of atomic models and averages.

**Example 10****Isothermal Titration Calorimetry**

NanR-binding duplex DNA (5'-gtttgaaaaaaatctcggtt atg-<sup>65</sup> gattattatggcgatggagattttcaaac-3', SEQ ID NO:15) was chemically synthesized. NanR was dialyzed extensively

against Tris buffer (50 mM Tris-HCl, pH 7.0; 300 mM NaCl) and the DNA was diluted to 0.03 mM in the same buffer. Protein and DNA samples were degassed by vacuum aspiration for 20 minutes prior to loading and titration was carried out at 25° C. The NanR dimer in the syringe (0.48 mM) was titrated against the 0.03 mM DNA sample in the reaction cell. Analysis of the interaction between the NanR/ManNAc-6P complex and the DNA sample was performed using 0.48 mM of complex and 0.03 mM of DNA. The calorimetric assays were performed using a VP-ITC (Micro-Cal Inc., Northampton, Mass.). The stirring speed was 300 rpm and the thermal power was recorded every 10 seconds. Data were processed and plotted using the Origin program (version 7) supplied with the instrument.

**Example 11****Sequence Comparison of NanR Protein**

<sup>50</sup> Conservation of *V. vulnificus*, *V. cholerae*, and *H. influenzae*-derived NanR sequences was examined. NanR amino acid sequences of *V. vulnificus* UJ016 (gi:37676858), *V. cholerae* O1 biovarE1 Tor str. N16961 (gi:15641775), and *H. influenzae* Rd KW20 (gi:16272110) were compared to each other. For sequence comparison, T-COFFEE software was used and for visualization, ESPript software was used. These two softwares are available in ExPASy portal (<http://au>>--<<expasy.org/>) (FIG. 13). High conservation was observed, and in particular, amino acids critical for ligand-

<sup>55</sup> binding and DNA-binding are conserved well.

**Experimental Example 1****Structure of NanR/ManNAc-6P Complex**

Unlike the typical structure of other transcriptional regulators, the NanR/ManNAc-6P complex analyzed by the

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above method contains two molecules in an asymmetric unit (A of FIG. 1). The symmetry mate in the dimer was analyzed, and a functional dimeric form of NanR was confirmed by electron microscopy (B of FIG. 1). The two NanR molecules at the dimer face each other and tilt approximately 45° in opposite directions (FIG. 2a). NanR adopts a two domain architecture that includes an N-terminal DNA-binding domain (DBD) and a large C-terminal ligand-binding domain (LBD) (FIG. 2b). The DBD comprises six helices while the LBD forms  $\alpha/\beta$  structure characterized by a five-stranded parallel  $\beta$ -sheet flanked by  $\alpha$  helices (FIG. 2b). The residues between  $\alpha$ 6 and  $\alpha$ 7 were not included in the final model because they were invisible in the electron density map; these regions are presumably very flexible. A DALI search revealed that no structures similar to that of NanR have been published. However, the structure of the LDB resembles the isomerase domain of glucosamine-6-phosphate synthase (GlmS), a bienzyme complex that catalyzes the first step in hexosamine metabolism. The fructose 6-phosphate binding site in the N-terminal isomerase subdomain of GlmS is located in the same position as that of the ManNAc-6P binding site in the NanR LBD, suggesting that NanR has evolutionarily adapted the isomerase domain to sense the nan regulatory molecule, whose structure is similar to that of fructose 6-phosphate. The structure of the NanR DBD is similar to the N-terminal domain of the *Bacillus subtilis* putative transcriptional regulator ybbH (PDB accession ID 2O3F). ManNAc-6P is located at the C-terminal edge of the  $\beta$  sheet in the LBD domain (FIG. 2b).

#### Experimental Example 2

##### Characterization of Interaction Between ManNAc-6P and NanR

An electron density difference map demonstrated that ManNAc-6P binds to NanR in the site formed by loops L9, L13, and L17 of the LBD domain (FIG. 3a). L13, which corresponds to the P-loop in the GlmS isomerase subdomain, crosses over and embraces the phosphate group of ManNAc-6P (FIGS. 3a and 3b); the phosphate oxygen atoms form hydrogen bonds with the side chains of S182, S184, and T187, and with the backbone amide of S183 in the P-loop (FIG. 3b, upper panel). This binding structure is very similar to that of the GlmS isomerase (FIG. 3b, lower panel). The side chain hydroxyl group of S138 in loop L9 is also hydrogen-bonded to a ManNAc-6P phosphate oxygen atom (FIGS. 3a and 3b). The hydroxyl group at position of the sugar ring forms a hydrogen bond with the A137 backbone amide. A hand-in-hand interaction between the two NanR monomers is formed by hydrogen bonds between the hydroxyl group at position O1 of the sugar ring in each NanR monomer and the nitrogen atom in the imidazole ring of each H163 (FIGS. 3a and 3c). This structure is critical for the conformational change of the NanR dimer and delivery of the signal to the nan operon genes when ligand-binding occurs. Furthermore, the phosphoryl group of the ligand forms a water-mediated hydrogen bond with the side chain amino group of R71 on  $\alpha$ 6 in the DBD (FIGS. 3a and 3c). These interactions may enable ligand-mediated relocation of the NanR dimer and influence its interaction with the nan operator. P231 and G234 form water-mediated hydrogen bonds with the carbonyl oxygen atom of the N-acetyl group (FIG. 3a). In addition, E229 and K240 form water-mediated hydrogen bonds with the sugar and phosphate oxygen atoms of ManNAc-6P (FIG. 3a).

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To assess the importance of the interaction between ManNAc-6P and NanR to regulation of the nan genes, the residues involved in ligand-binding were mutated and the impact on NanR function was investigated using an *E. coli* dual plasmid system. Cells were cotransformed with plasmids containing a luciferase reporter gene fused to the NanR-binding nanT(PSL)AR promoter (PnanTp), and wild-type or mutant NanR, and then incubated in the presence of arabinose and the presence or absence of Neu5Ac. Luciferase activity in cells expressing the wild-type NanR was increased following addition of Neu5Ac; however, Neu5Ac was unable to activate PnanTp in cells expressing the mutant NanR (S138A, H163A, H163L, S182A, E229L, K240A or K240M), except the S184A and T187A mutants (FIG. 3d). These results suggest that precise binding of ManNAc-6P, the metabolic intermediate of Neu5Ac to NanR is critical for regulation of the nan genes.

#### Experimental Example 3

##### Characterization of DBD of NanR

The simplest helix-turn-helix (HTH)-containing DBD includes three core helices; the HTH domains can form tetra-helical bundle, winged helix, and ribbon-helix-helix type configurations. The DBD of NanR is composed of a six-helix bundle, which is not the archetypal conformation of HTH-containing domains; therefore, the recognition helix required for DNA-binding could not be determined. Nevertheless, analysis of the surface electrostatic potential of NanR revealed a number of positively charged residues in the DBD domain (K20, K21, R23, R57, R60, and K65) that may be responsible for binding to the phosphate backbone of DNA (FIGS. 4a and 4b). The importance of these residues to DNA-binding was examined using an *E. coli* dual plasmid system described in Example 2. The repressive effect of wild-type NanR on activity of PnanTp was abolished by the R57A, R57L, R60A, and R60L point mutations (FIG. 4c). Immunoblot analyses using anti-NanR antiserum revealed that these results were attributable to functional defects of the mutants and not to reduced cellular expression (FIG. 5).

NanR represses both nan T(PSL)AR and nanEK nagA operons by binding to an operator within the nanTp-nanE intergenic region. To examine the role of R57 and R60 residues in DNA binding, EMSA (electrophoretic gel mobility shift assays), in which the nanTp-nanE intergenic region was incubated with wild-type or mutant NanR in the absence or presence of ManNAc-6P, were performed. In the experiment using the wild-type NanR, addition of ManNAc-6P resulted the retardation of DNA migration, suggesting that ManNAc-6P alters the ability of NanR to bind to the nan operator. No DNA migration was detected in the experiment using the mutant NanR(R57A or R60A) (FIG. 4d). With the exception of the S184A mutant, which was as active as wild-type NanR in the *E. coli* dual plasmid system experiments, the DNA migration by the ligand-binding defective mutants of NanR was not affected by addition of ManNAc-6P (FIG. 4d).

In addition, an in vitro transcription assay revealed that the R57A and R60A mutants did not repress PnanE even in the absence of ManNAc-6P, unlike the wild-type and S184A mutant (FIG. 4e). Consistent with the results of EMSA and the *E. coli* dual plasmid system assay, the other ligand-binding defective mutants of NanR did not enable transcription of nanE in either the presence or absence of ManNAc-6P (FIG. 4e). These results indicate that R57 and R60 in  $\alpha$ 5 are indispensable for binding of NanR to the nan operator.

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and that the DNA-binding HTH motif in each NanR monomer includes  $\alpha$ 4 and  $\alpha$ 5. The distance between the two  $\alpha$  helices in the NanR dimer is approximately 22 Å (FIG. 4b), suggesting that the method of DNA-binding employed by NanR differs from that of other HTH motif containing transcriptional regulators.

#### Experimental Example 4

##### ManNAc-6P Alleviates the Interaction Between NanR and Transcriptional Control Region of Nan Operon

The EMSA and in vitro transcription experiments demonstrated that S138, H163, S182, E229, and K240 residues are critical for ligand-sensing and regulation of the nan genes by NanR. Therefore, the present inventors hypothesized that binding of ManNAc-6P may alter the conformation of NanR and alleviate its interaction with the transcriptional control region of nan operon. In detail, a DNaseI footprinting assay using NanR protein and  $^{32}$ P-labeled transcriptional control region of nan operon was performed. Upon addition of ManNAc-6P, cleavage bands at the center of the NanR-binding site disappeared and neighboring regions were de-protected by the protein (FIG. 6). Isothermal titration calorimetry analyses revealed a robust interaction between the NanR dimer and the target DNA, with a 1:1 binding stoichiometry and a dissociation constant ( $K_d$ ) of 1.40  $\mu$ M. However, the interaction affinity of NanR complexed with ManNAc-6P was reduced by 130-fold ( $K_d=185.87 \mu$ M). Taken together, these results support the hypothesis that ManNAc-6P alters the conformation of NanR via relocation of ligand-binding residues, thereby reducing nan-binding activity of NanR which is a transcriptional repressor of the nan genes.

#### Experimental Example 5

##### Electron Microscopic Analysis of the Interaction Between NanR and Nan Genes

Electron microscopy using negative staining followed by single particle analysis showed that apo-NanR and the NanR/ManNAc-6P complex share similar structural features when analyzed at the molecular level of approximately 2 nm resolution (FIG. 7a; A and B of FIG. 9). Additional electron microscopy densities were observed for the DNA-bound NanR dimer (FIGS. 7a and 7b, indicated by white arrows; FIG. 5C). The DNA binding pattern revealed that it passes between the DBD domains in the NanR dimer.

It was confirmed that  $\alpha$ 5 of each NanR molecule is essential for DNA-binding. The distance between the  $\alpha$ 5 helices that project towards the interior of the DBD domains is approximately 22 Å (FIG. 4b), which is close to the width of the DNA double helix (20 Å). Two-dimensional fitting demonstrated that the atomic models assembled from the crystal structures of NanR and DNA (FIG. 7b, middle panel) fit onto the averaged image of the NanR/DNA complex well (FIG. 7b, right panel). However, the entire length of DNA associated with NanR does not match perfectly due to its flexibility.

Meanwhile, inspection of the surface electrostatic potential of all in the LBD of NanR revealed the existence of positively charged residues that may also be responsible for DNA-binding (FIG. 7C). The importance of K188 and K199 residues on all to binding of NanR to DNA was assessed by EMSA. Although the K188A, K188L, K199A, and K199L

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NanR mutants were able to bind to DNA, the binding activity of the mutants was not as conspicuous as that of wild-type NanR (FIG. 10). Furthermore, the mutants were much more susceptible to ManNAc-6P than wild-type NanR (FIG. 10). Taken together, these results demonstrate that the NanR dimer forms an arched tunnel-like DNA-binding space that is formed mainly by  $\alpha$ 5 and  $\alpha$ 11 in each monomer. The transcriptional control region of nan operon interacts with the dimer via the positively charged residues in this space (FIGS. 7b and 7c).

#### Experimental Example 6

##### Effect of Interaction Between ManNAc-6P and NanR on Growth and Survival During Infection

Robust control of the genes encoding catabolic enzymes and the putative transporter for Neu5Ac is crucial for growth and survival of pathogenic bacteria in the host. To investigate the biological relevance of ligand-sensing by NanR, the present inventors examined the effects of mutation of R57 and H163, which are critical for DNA-binding and ligand-binding, respectively, on growth of the pathogenic bacteria *V. vulnificus*. The R57A or H163L mutation was introduced into *V. vulnificus* chromosomal DNA and the in vitro growth of each mutant strain was examined. Growth of the R57A strain was similar to that of the wild-type strain; however, growth of the H163L strain was impaired in minimal M9 medium supplemented with Neu5Ac as a sole carbon source (FIG. 11a). Addition of D-xylose and L-proline restored the growth of the H163L strain (which had an altered colony morphology with reduced opacity), as observed previously for the nanA mutant strain. In the supplemented medium with D-xylose, L-proline and Neu5Ac, the expression levels of nan genes in the H163L nanR mutant were at least 142-fold lower than those in wild-type nanR (FIG. 12). These results suggest that the H163L mutant is indeed defective for ManNAc-6P sensing and nan gene regulation, thus affecting the growth of *V. vulnificus* when Neu5Ac is available. To examine the importance of the residue for ligand-binding, a mouse intestine colonization competition assay was performed. In eight of the 10 mice studied, colonization of the H163L mutant strain was 22.4-fold lower than that of the wild-type strain, resulting in a median competitive index of 0.045 (FIG. 11b).

Finally, mice were challenged with a lethal dose of *V. vulnificus*. At 24 hours after infection, the percentages of surviving mice challenged with *V. vulnificus* expressing the H163L mutant or wild-type strain were 60% and 20%, respectively (FIG. 11c). These results indicate that regulation of NanR by ManNAc-6P is required not only for growth and survival, but also for the pathogenesis of *V. vulnificus*.

Taken together, the NanR protein is an important protein that affects growth, survival, and pathogenesis of a variety of bacteria having nan genes, since it binds to nan operator to suppress nan operon expression. It was confirmed that the functions of NanR are regulated by binding of its regulatory ligand, ManNAc-6P. Therefore, substances or new drugs against symptoms caused by a variety of bacteria having nan genes can be developed by regulation of nan gene expression using the three-dimensional structure of the NanR/ManNAc-6P complex of the present invention.

Based on the above description, it will be apparent to those skilled in the art that various modifications and changes may be made without departing from the scope and spirit of the invention. Therefore, it should be understood that the above embodiment is not limitative, but illustrative in all aspects. The scope of the invention is defined by the appended claims rather than by the description preceding them, and therefore all changes and modifications that fall within metes and bounds of the claims, or equivalents of

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such metes and bounds are therefore intended to be  
embraced by the claims.

## EFFECT OF THE INVENTION

The NanR protein is an important protein that affects growth, survival, and pathogenesis of a variety of bacteria.

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having nan genes, since it binds to nan operator to suppress nan operon expression. Therefore, new drugs for the prevention or treatment of diseases associated with a variety of bacteria having nan genes can be developed using a crystal 5 of NanR and its regulatory ligand ManNAc-6P complex, a crystallization method, and a three-dimensional structure thereof.

**SEQUENCE LISTING**

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<212> TYPE: PRT

<213> ORGANISM: *Vibrio vulnificus*

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                  20                 25                 30

Ala His Asp Val Gln Phe Gln Thr Ile Thr Asp Leu Ala Arg Asn Thr  
35 40 45

Gln Thr Ser Glu Ala Thr Val Val Arg Leu Cys Arg Asp Met Gly Tyr  
 50                    55                    60

Lys Gly Tyr Ser Asp Phe Arg Met Ala Leu Ala Val Asp Leu Ser Gln  
65                   70                   75                   80

Thr Glu Ser Arg Gln Gln Asn His Ile Glu Gly Asp Ile Cys Asp Val  
85 90 95

Ser Ala Gln Ser Ala Val Asp Ser Leu Gln Asp Thr Ala Lys Leu Ile  
100 105 110

115                    120                    125

Phe Ile Gly Cys Ile Gly Val Gly Ala Ser Ser Ile Val Gly Arg Tyr

Leu Ala Tyr Arg Leu Ile Arg Ile Gly Lys Lys Ala Ile Met Phe Glu

Asp Thr His Leu Ala Ala Met Ser Ala Ser Arg Ser Ser Gln Gly Asp  
165 170 175

Leu Trp Phe Ala Val Ser Ser Ser Gly Ser Thr Lys Glu Val Ile His

Ala Ala Gly Leu Ala Tyr Lys Arg Asp Ile Pro Val Val Ser Leu Thr

Asn Ile Asn His Ser Pro Leu Ser Ser Leu Ser Thr Glu Met Leu Val  
GAG GAT GAT GTC TGT TGT

Ala Ala Arg Pro Glu Gly Pro Leu Thr Gly Gly Ala Phe Ala Ser Lys  
225 226 227 228 229 230 231 232 233 234 235 236 237 238

Val Gly Ala Leu Leu Leu Val Asp Val Leu Val Asn Ser Leu Leu Glu

Ser Tyr Pro Glu Tyr Lys Asp Ser Val Gln Glu Thr Ala Glu Val Val

Ile Pro Leu Met Ala Asn

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<210> ORGANISM: Vibrio vulnificus  
<211> LENGTH: 837  
<212> TYPE: DNA
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<210> SEQ ID NO: 2  
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<212> TYPE: DNA  
<213> ORGANISM: *Vibrio vulnificus*

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-continued

&lt;400&gt; SEQUENCE: 2

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aaaactgcgtg ttgtggcgga ctacatattg gaaaatgcgc atgatgtgca gtttcaaacc      120
atcacggatc ttgctcgcaa cacacaacc agtgaagcga cagtcgtacg cttatgtcgc      180
gacatgggct ataagggcta ttccgattt cgtatggcgc ttgccgttga tttgagccaa      240
accgaaagtc gtcagaaaa tcatacgaa ggtgacattt gcatgtgtc tgccaaagc      300
gggttagaca gcctgcaaga caccgcaaaa cttatcgatc gttaatcatt ggccgcatt      360
gttgagcggg ttcatcaagc cgagtttattt ggggtcattt gtgttggagc gtcgagcatt      420
gttggccgtt acctcgctta tcgccttata cgtatcggtt agaaagcgat catgtttgaa      480
gataccatt tagccgcaat gagtgcaagc cgctcaagtc aaggatgtatct gtggtttgc      540
gttccagtt cagggttcgac gaaagaagtc attcatgccc ctgggttgc gtataaggcgt      600
gatattcccg tcgtttctct gacaaacatc aatcacagcc cgctctcttc tctctcaact      660
gaaatgtgg tggctgcaag accagaaggg ccactaacag gtggtgctt tgccctgaaa      720
gtcgccgcgc tgctcttagt ggacgtgtt gtcaacttattt tagagag ctacccggaa      780
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&lt;211&gt; LENGTH: 299

&lt;212&gt; TYPE: PRT

&lt;213&gt; ORGANISM: Vibrio vulnificus

&lt;400&gt; SEQUENCE: 3

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Asp	Ser	Asn	Agn	Gln	Val	Asn	Tyr	Ala	Val	Ile	Asp	Gln	Ile	Ala	Ala
20															

Leu	Leu	Ile	Glu	Gln	Gly	Val	Thr	Gly	Ala	Tyr	Val	Cys	Gly	Thr	Thr
35															

Gly	Glu	Gly	Ile	His	Cys	Ser	Val	Glu	Glu	Arg	Lys	Ala	Ile	Ala	Glu
50															

Arg	Trp	Val	Lys	Ala	Ala	Asp	Gly	Lys	Leu	Asp	Ile	Ile	Leu	His	Thr
65															

Gly	Ala	Leu	Ser	Ile	Val	Asp	Thr	Leu	Glu	Leu	Thr	Arg	His	Ala	Glu
85															

Thr	Leu	Asp	Ile	Leu	Ala	Thr	Ser	Ala	Ile	Gly	Pro	Cys	Phe	Phe	Lys
100															

Pro	Ser	Ser	Val	Ala	Asp	Leu	Val	Asn	Tyr	Cys	Ala	Gln	Ile	Ala	Glu
115															

Ala	Ala	Pro	Ser	Lys	Gly	Phe	Tyr	Tyr	Tyr	His	Ser	Gly	Met	Ser	Gly
130															

Val	Asn	Leu	Asp	Leu	Glu	Gln	Phe	Leu	Ile	Gln	Gly	Glu	Gln	Arg	Ile
145															

Pro	Asn	Leu	Ser	Gly	Ala	Lys	Phe	Asn	Asn	Val	Asp	Leu	Tyr	Glu	Tyr
165															

Gln	Arg	Ala	Leu	Arg	Val	Ala	Asn	Gly	Lys	Phe	Asp	Ile	Pro	Phe	Gly
180															

Val	Asp	Glu	Phe	Leu	Pro	Ala	Gly	Leu	Ala	Val	Gly	Ala	Ile	Gly	Ala
195															

Val	Gly	Ser	Thr	Tyr	Asn	Tyr	Ala	Ala	Pro	Leu	Tyr	Leu	Lys	Ile	Ile
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**479****480**

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210            215            220

Glu Ala Phe Asn Gln Gly Lys His Ser Glu Val Gln Ala Leu Met Asp  
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 Lys Val Ile Ala Leu Ile Arg Val Leu Val Glu Tyr Gly Val Ala  
 245                250                255  
 Ala Gly Lys Ala Ala Met Gln Leu His Gly Ile Asp Ala Gly Asp Pro  
 260                265                270  
 Arg Leu Pro Ile Arg Ala Leu Thr Ala Gln Gln Lys Ala Asp Val Val  
 275                280                285  
 Ala Lys Met Arg Asp Ala Asp Phe Leu Asn Leu  
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<210> SEQ ID NO 4  
 <211> LENGTH: 427  
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 <213> ORGANISM: Vibrio vulnificus

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 Pro Val Gly Phe Ser Leu Ile Phe Val Ala Leu Val Phe Leu Ile Val  
 20                25                30  
 Thr Asn Ser Thr Gly Ile Asn Phe Ala Ala Gln Gln Met Leu Gly Gly  
 35                40                45  
 Ile Asp Asn Phe Thr Leu Leu Ala Val Pro Phe Phe Val Leu Thr Gly  
 50                55                60  
 His Leu Met Asn Ser Ala Gly Ile Thr Glu Arg Ile Phe Asn Phe Ala  
 65                70                75                80  
 Lys Ser Met Val Gly His Ile Thr Gly Ser Leu Gly His Val Asn Ile  
 85                90                95  
 Met Ala Ser Leu Leu Phe Ser Gly Met Ser Gly Ser Ala Leu Ala Asp  
 100                105                110  
 Ala Gly Gly Leu Gly Gln Leu Glu Ile Lys Ser Met Arg Asp Ala Lys  
 115                120                125  
 Tyr Asp Asp Asp Phe Ala Gly Gly Leu Thr Ala Ala Ser Cys Ile Ile  
 130                135                140  
 Gly Pro Leu Val Pro Pro Ser Val Pro Leu Val Ile Tyr Gly Val Val  
 145                150                155                160  
 Ser Asn Thr Ser Ile Gly Ala Leu Phe Leu Ala Gly Ala Ile Pro Gly  
 165                170                175  
 Ile Leu Cys Cys Val Ala Leu Met Val Met Ser Tyr Phe Ile Cys Lys  
 180                185                190  
 Lys Arg Gly Tyr Met Thr Leu Pro Lys Ala Ser Arg Lys Glu Gln Phe  
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 Thr Ser Phe Lys Glu Ala Phe Leu Ser Leu Met Thr Pro Val Ile Ile  
 210                215                220  
 Ile Gly Gly Ile Phe Ser Gly Lys Phe Thr Pro Thr Glu Ala Ala Val  
 225                230                235                240  
 Val Ser Ser Leu Tyr Ala Leu Phe Leu Gly Thr Val Val Tyr Lys Gln  
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 Leu Thr Leu Thr Gly Phe Val Glu Ile Leu Arg Glu Thr Val Asn Thr  
 260                265                270  
 Thr Ala Val Val Ala Leu Met Val Met Gly Val Thr Val Phe Gly Trp  
 275                280                285

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-continued

Ile Val Ala Arg Glu Gln Leu Pro Gln Met Leu Ala Asp Tyr Phe Leu  
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Ser Ile Ser Glu Asn Pro Leu Val Leu Leu Leu Ile Asn Leu Leu  
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Leu Leu Phe Leu Gly Thr Phe Ile Glu Ser Leu Ala Leu Leu Leu  
 325 330 335

Leu Val Pro Phe Leu Val Pro Val Ala Ser Ala Val Gly Ile Asp Pro  
 340 345 350

Val His Phe Gly Val Met Ala Ile Leu Asn Leu Met Ile Gly Ile Leu  
 355 360 365

Thr Pro Pro Met Gly Met Ala Leu Tyr Val Val Ser Arg Val Gly Asp  
 370 375 380

Ile Pro Phe His Thr Leu Thr Arg Gly Val Leu Pro Leu Leu Val Pro  
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Leu Phe Ile Val Leu Ala Leu Val Ala Val Phe Pro Gln Ile Thr Leu  
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Leu Leu Pro Glu Leu Leu Leu Gly Tyr Gly Gln  
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<210> SEQ ID NO 5  
 <211> LENGTH: 168  
 <212> TYPE: PRT  
 <213> ORGANISM: Vibrio vulnificus

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Trp Leu Leu Asn Asp Pro Ser Leu Trp Ser Glu Glu Leu Ala Arg Val  
 35 40 45

Leu Phe Met Tyr Met Ser Leu Ile Gly Cys Ala Ile Ala Ile Lys Arg  
 50 55 60

Gly Thr His Val Asn Ile Thr Phe Phe Ser Asp Lys Leu Pro Glu Lys  
 65 70 75 80

Ile Arg Leu Leu Val Leu Ser Leu Glu Ala Ala Val Leu Val Ser  
 85 90 95

Ile Phe Ala Ile Ile Tyr Leu Gly Tyr Gln His Val Glu Arg Thr Ala  
 100 105 110

Phe Phe Glu Leu Ile Thr Leu Gly Val Ser Ser Lys Trp Met Asn Tyr  
 115 120 125

Ser Leu Pro Leu Gly Gly Leu Phe Met Val Ile Arg Gln Leu Gln Lys  
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 <211> LENGTH: 321  
 <212> TYPE: PRT  
 <213> ORGANISM: Vibrio vulnificus

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-continued

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Ser Val Gly Ser Val Glu Tyr Thr Ser Ala Lys Leu Leu Ala Asp Thr  
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Val Glu Glu Met Ser Lys Gly Glu Leu Lys Leu Ala Leu Tyr Pro Ser  
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Ala Gln Leu Gly Asp Asp Arg Ala Met Leu Gln Gln Leu Ser Met Gly  
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Pro Arg Ala Glu Ala Val Thr Leu Pro Tyr Val Ala Arg Asp Tyr Asp  
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His Leu Arg Arg Met Phe Asp Ser Glu Phe Gly Gln Gly Ile Arg Gln  
115 120 125

Glu Met Leu Thr Lys Phe Asn Trp Arg Ala Leu Asp Thr Trp Tyr Asn  
130 135 140

Gly Thr Arg Glu Thr Thr Ser Asn Arg Pro Leu Lys Ser Ile Ser Asp  
145 150 155 160

Phe Lys Gly Leu Lys Leu Arg Val Pro Asn Ala Lys Pro Asn Leu Asn  
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Tyr Ala Lys Leu Ser Gly Ala Ser Pro Thr Pro Met Ala Phe Ser Glu  
180 185 190

Val Tyr Leu Ala Leu Gln Thr Asn Ala Val Asp Gly Gln Glu Asn Pro  
195 200 205

Leu Pro Thr Ile Lys Thr Met Lys Phe Tyr Glu Val Gln Ser Asn Leu  
210 215 220

Ala Ile Thr Asn His Ile Val Asn Asp Gln Met Val Leu Ile Ser Glu  
225 230 235 240

Ser Thr Trp Gln Lys Leu Ser Glu Gln Glu Arg Glu Ile Val Ala Asn  
245 250 255

Ala Val Lys Gln Thr Gly Glu Ala His Thr Ala Ser Val Lys Lys Gln  
260 265 270

Glu Ala Glu Leu Ile Ser Phe Phe Glu Ala Gln Gly Val Asn Val Thr  
275 280 285

Tyr Pro Glu Leu Ala Pro Phe Arg Glu Ala Met Gln Pro Leu Tyr Ser  
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Glu Phe Glu Lys Lys Ile Gly Gln Pro Ile Val Ser Lys Leu Ala Ala  
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Met

&lt;210&gt; SEQ ID NO 7

&lt;211&gt; LENGTH: 225

&lt;212&gt; TYPE: PRT

&lt;213&gt; ORGANISM: Vibrio vulnificus

&lt;400&gt; SEQUENCE: 7

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20 25 30

Gln Ala Gly Ala Lys Ala Leu Arg Ile Glu Gly Val Glu Asn Val Arg  
35 40 45

His Val Ser Gln Ala Thr Asn Val Pro Ile Ile Gly Ile Val Lys Arg  
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**485****486**

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					115			120					125		
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					130			135					140		
Gly Tyr Val Gly Glu Ile Glu Pro Thr Glu Pro Asp Leu Glu Leu Val															
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Lys Gln Phe Ser Ser Ala Gly Phe Phe Thr Met Ala Glu Gly Arg Tyr															
					165			170					175		
Asn Thr Pro Gln Leu Ala Ala Lys Ala Ile Glu Asn Gly Ala Val Ala															
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Val Thr Val Gly Ser Ala Ile Thr Arg Met Glu Val Val Thr His Trp															
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Tyr															
					225										

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						20			25			30			
Val Val Asn Asp Ala Thr Thr Leu Ala Lys Glu Ile Leu Ala His Cys															
						35			40			45			
Gln Ala Trp Leu Ser Asp Val Asp Ala Ile Gly Ile Ser Thr Thr Gly															
						50			55			60			
Leu Val Ser Glu Gln Gly Ile Ser Ala Ile Asn Pro Gly Thr Leu Ser															
						65			70			75			80
Phe Pro Thr Pro Phe Pro Leu His Ser Glu Leu His Arg Leu Ser Gly															
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Lys Pro Val Lys Met Leu Asn Asp Ala Gln Ala Ala Ala Trp Tyr Glu															
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Phe Leu Gln Leu Ser Pro Glu Leu Asp Val Arg Asn Met Ala Tyr Ile															
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Thr Val Ser Thr Gly Val Gly Gly Leu Val Ile Asn Gln Gln Leu															
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His Lys Gly Lys Ser Asn Phe Ala Gly His Ile Gly His Thr Val Leu															
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Asp Pro Asn Gly Pro Leu Cys Gly Cys Gln Gln Arg Gly Cys Val Glu															
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Ala Ile Ala Ser Gly Asn Ala Ile Asn Ala Gly Ala Gln Ala Leu Phe															
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Glu Gln Ala Ser Ala Leu Ile Gln Gln Ser Ala Glu Ala Ile Ala Gln  
 210 215 220

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 225 230 235 240

Gly Gly Gly Val Gly Leu Ala His Gly Tyr Leu Ala Arg Val Gln Ala  
 245 250 255

Phe Ile Asp Lys Gln Pro Leu Val Phe Gln Val Lys Val Arg Ala Ala  
 260 265 270

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&lt;213&gt; ORGANISM: Vibrio vulnificus

&lt;400&gt; SEQUENCE: 9

atgatgaaca aattaaaagg gctgatcgcc gtcctcata caccgttga tagcaataat 60  
 caggtgaact atgcggtgat cgatcaaatt gctgcattac tcattgaaca aggtgttaca 120  
 ggggcttacg tatgtggcac cacccggaa ggcatacatt gttcagttga agagegaaaa 180  
 gcatgttgcg agcgttgggt gaaagccgca gatggtaagt tagacattat tcttcatacg 240  
 ggggctctga gtatcggtga taccctttag cttacacgac acgctgaaac actggatatt 300  
 ctggccactt ccgcgatcg gccttgcattt ttcaaaaccga gcagcgtcgc tgatcttgc 360  
 aattactgcg cgcatgttgc ggaagccgca ccgtcaaaag gtttcttatta ctaccactca 420  
 ggaatgtctg gcgtgaatct cgatctggaa cagttctca tccaaggaga gcagegtatt 480  
 ccgaacttat ctggggccaa gtttataac gtggatctt atgagttacca acgcgcgctg 540  
 agagtagcca atggcaagtt tgatattca ttccggcggtt atgaattttt gccagcagg 600  
 ctggccctgtt gggctattgg ggcgggttgc agcacttata attacgccgc accactgtat 660  
 ctgaagatca ttgaggcggtt caaccaaggg aaacatagtg aagttcaggc gttgatggat 720  
 aaagtgtatcg cattgtatcg tttttttttt ggatccgggtt ggcggcggtt agggaaagca 780  
 gcgatgcaat tacacggcat tgatgccgtt gatccgcgtt tgccaaattcg acgcgtcact 840  
 gcgcaacaaa aagccgatgt cgtggcgaaa atgcgcgtt cggatccat taatctgtat 900

&lt;210&gt; SEQ ID NO 10

&lt;211&gt; LENGTH: 1284

&lt;212&gt; TYPE: DNA

&lt;213&gt; ORGANISM: Vibrio vulnificus

&lt;400&gt; SEQUENCE: 10

atggctgggtt caatttttgg ttggcttaggc ttgttttttgc ctggcatgcc agttggcttt 60  
 tctctgattt ttgtggcggtt ggttttttttt attgttacca acagcacagg tattttttttt 120  
 gcaagcgcagc agatgtgtggg cggcatcgat aattttacgt tattggcggtt accattttttt 180  
 gtccttaacgg gccattttgtat gaacagtgcg ggttattacgg agcgaattttt taactttgc 240  
 aaatccatgg ttggccatat cacgggcagt cttagggcatg taaacatcat ggcgagttta 300  
 ctctttctgtt gatgtctgg ctctgcgtt gccgatgcgg gaggcttggg gcagcttgc 360  
 atcaaatcaa tgcgtgacgc gaaatacgat gatgatccat ctgggtggctt cacggcgcc 420

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tctgtccatca ttggtccgct tggccgcct tctgtccat tggtgattta tggtgtggc	480
tccaaacatcgatggcgtc gttgtttta gcccgtgc aaaccaggcat attgtgcgc	540
gtggcttgc ttgtgtatgc ctatttcata tgtaagaaac gtggctacat gacccgtc	600
aaaggcgtcga gaaaggagca gttcacatcg ttcaaaaggagg catttttc tctaattact	660
cctggcatca tcattggcgg gatctttcg ggtaagtca ccccgacggc agcagcgtc	720
gttccctctc ttacgcttt gttcccttgc accgtgggtc acaaacacgt gacccgtc	780
gggtttgtgg aaattctgcg cgaaaccgtg aataccaccg ctgttgc tttgtatggc	840
atggggtaa cgggtttgg ctggatcgat gcgcgtgc aattaccaca gatgttgc	900
gattatttct ttgtcgtatcg taaaaccca ttggactacat tgctactcat caacttgc	960
ttgtctgtttt tggaaacctt cattgaatca ctgcgcgtat tgctattgtc gttccattt	1020
tttagccccag tggcatcagc ggtcgggatt gaccctgtgc actttgggt tatggccatt	1080
ttgaacttgc tgatcggcat tctcacgcca ccaatggcata tggcttata tgggtgtca	1140
cgtgtcggag acattccatt tcataccttactcgaggcg tacttccctct tctggccca	1200
ttgtttattt tgctggcgct ggtggcgta ttccctcaaa tcaccttgc attgcctgag	1260
ctgttttgg gttacggaca ataa	1284

&lt;210&gt; SEQ ID NO 11

&lt;211&gt; LENGTH: 507

&lt;212&gt; TYPE: DNA

&lt;213&gt; ORGANISM: Vibrio vulnificus

&lt;400&gt; SEQUENCE: 11

atgttacgca agatcttcga taattttagaa gagatcatca cagtgcact tatggccctct	60
cttcttggc tgctgacatcg ccaaatttgcg acgcgcgtgg tacttaacgc tccttacta	120
tggagtgaag agttacgcgc agtgcgtttt atgtatatgt cgctcattgg ctgtgcac	180
gctatcaaactca gttggactca cgtcaatatac acgttttctt cagataagtt gccagaaaa	240
atacgccctt tgggtgtttt gtccttggaa gctgcgttat tggtttgcattt attcgccatc	300
atttatttgg gctatcaaca cgtggaaaga accgcctttt ttgaattgtat taccttaggt	360
gtgtccagca aatggatgaa ctacagtttta ccccttggcg ggctttcat ggtgattcgt	420
caattgcaaa aaatggtcgg tattgtgcgcg gaatttcgtc acgagtgcgg tgcgttacc	480
gcgtcgaatc atgcagagca aaggtaa	507

&lt;210&gt; SEQ ID NO 12

&lt;211&gt; LENGTH: 966

&lt;212&gt; TYPE: DNA

&lt;213&gt; ORGANISM: Vibrio vulnificus

&lt;400&gt; SEQUENCE: 12

atgaaaacca tgaaccgaat tacacttgcgt atgctgactg tgggatttgcg ctttcttcg	60
aatgctgcaa cgacgttgcgaa aatggaaatgc caagcgtcggttggctcagt ggaatacacc	120
tctgccaagc tgcttgcgcg cacgggttgcgaa gagatgtatgc aggggtgatgtt gaaatcgatcg	180
ttatatccaa gtgcgcact gggcgacgcg cgtgcgtatgt tacagcgtcttctatgggt	240
gacttagaca tcacttacgc ggaatttggc cgtatggac tttggatccc aagagcagaa	300
gggggtgcacct taccttacgt tgctcgtatgtatgaccatt tacgcgcgtat gtttgcattcc	360
gagtttggcc aaggcattcg tcaggaaatgc ctaaccaatgt ttaattggcg tgctgtatgt	420
acttggtaca acggaaacacgc tggaaaccacc tctaaccgtc cattgtatgtc aatttcgtatgt	480

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tttaaaggc tgaaattgcg cgtgccgaat gcgaaaccaa atctgaatta cgccaagttg	540
tctggggcat caccaacacc gatggcatt tctgaagttt acctcgcgct acaaaccaat	600
gcagttgacg ggcaagagaa cccattgcca accattaaga caatgaagtt ctacgaggtg	660
caaagcaact tggcgatcac caatcacatc gtgaacgacc agatggtgct gatttcagaa	720
agcacgtggc aaaagctctc tgagcaggaa cgtgaaattt tcgctaacgc ggtgaaacaa	780
acaggggaaag cgcataccgc atcgggtaaa aagcaagaag cggagctgat ctcttcttt	840
gaggegcagg gtgtgaatgt cacctaccca gaacttgcgc cgttccgtga agccatgcag	900
ccgccttact cagagttga aaagaaaatt ggtcagccga tcgtgtctaa gttggcggca	960
atgttag	966

&lt;210&gt; SEQ ID NO 13

&lt;211&gt; LENGTH: 678

&lt;212&gt; TYPE: DNA

&lt;213&gt; ORGANISM: Vibrio vulnificus

&lt;400&gt; SEQUENCE: 13

atgccttcgg gccagactgt tgtttcaatt cagcctgttg tcggtagccc tcttgataaa	60
accgagttca tcgttctat ggcagttgt gcagaacaag coggagotaa ggcactgcgt	120
attgaaggcg ttgaaaacgt acgcctatgt tctcaagcga ccaatgtacc aataatttggaa	180
attgtgaaac gtgatttaca ggacagtctt gtgcgcacca caccttcgt ctgtgacgtg	240
gtatgcctcg cgactgctgg cgcaaccatc atgcgccttg atgcgaccga tcgtcagcga	300
ccagagagcc gagaaacgtat agccaacgcata atcaaaaaca gtgggtgtt cgcgatggca	360
gattgcctt gcttcgcggta tggcaatgg gcccccaaa ttggggtcga tattattggc	420
tcaaccttgc ctggctacgt gggcgaaattt gaaccaactg aacccgattt ggaactggtg	480
aaacagttt ctccggcgaa atttttacc atggcgaaat gtcgctacaa cacgcctcaa	540
ctcgccgcca aagcgattga aaatggcgat gtcgcccgtga cggggcgttc ggcacatcaca	600
cgaatggaaatg tggttactca ctggtttaat tctgcaacac aagcggtaaatg acagaataat	660
gaaagtattttaa gctatttgc	678

&lt;210&gt; SEQ ID NO 14

&lt;211&gt; LENGTH: 882

&lt;212&gt; TYPE: DNA

&lt;213&gt; ORGANISM: Vibrio vulnificus

&lt;400&gt; SEQUENCE: 14

atgaaaggat tagctatttgc tattggcggt aaaaaatcg ctttaggcaa cgtggtcgtat	60
gggcatttgc agcatcgcaaa acatggcatttgc acccctgttg tcaacgcgtgc aacaacccaa	120
gcgaaagaga ttcttgccttgc ctgccaagca tggcttaagcg acgttgcgtgc catcggtatt	180
tgcactaccgc ggcttgcgtat cgaacaaggaa ataagtgccaa ttaatccagg cacattggac	240
ttccctacgc cttttctctgc gcatagcgaa ctacacagat taagtggccaa gccagttaaa	300
atgctcaatgc atgctcaatgc ggctgcctgg tatgagttt tgcacacttgc acctgactg	360
gatgttcgca atatggccta tattactgtc tcgacaggtt tggggaggcggtt tttgggtatt	420
aaccaacaac tgcataaggaa taagtcaaat tttggccggat atattggccaa tacgggttt	480
gatccaaatgc gcccacttttgc tggctgccttgc cagcgaggat ttgttgcgttgc catcgcttgc	540
ggaaatgccttgc ttaacgcagg cgctcaagcc ctttttggccaa aacgcatttc taatatcgaa	600

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ctgtttcagc ttgctcaaca caatgaacaa gcctccgcac tcattcaaca aagcgccgaa	660
gcctatgcggc agctctgcct aaataaaa gccacgctgg atctcgattt ggtggtcata	720
ggcggeggtg tgggccttgc tcacggctac ctgcgtcgta tacaaggcgtt tatcgacaaa	780
cagcctctcg tgtttcaggt taaggtgaga gcccagttt gcatgatcgatcgatcgatc	840
cttggcgcag ctttcaatt tgaggagat aatcttcattt ga	882

<210> SEQ ID NO 15  
<211> LENGTH: 110  
<212> TYPE: DNA  
<213> ORGANISM: Artificial Sequence  
<220> FEATURE:  
<223> OTHER INFORMATION: Synthetic NanR-binding duplex DNA

<400> SEQUENCE: 15

gtttgaaaaa aatcttcgtt atggattattt atggcgatgg agattatttt caaaccaaac	60
tttttttaga agcaataacctt aataataccg ctacctctaa taaaactttt	110

<210> SEQ ID NO 16  
<211> LENGTH: 278  
<212> TYPE: PRT  
<213> ORGANISM: Vibrio cholerae

<400> SEQUENCE: 16

Met Asn Leu Pro Lys Asn Leu Met Val Arg Leu Arg Ser Asn Thr Arg	
1 5 10 15	
Pro Ile Ser Lys Lys Leu Arg Val Val Ala Asp Tyr Val Leu Phe Asn	
20 25 30	
Ala His Arg Val Gln Tyr Gln Thr Ile Thr Asp Leu Ala Pro Asn Thr	
35 40 45	
Lys Thr Ser Arg Ala Thr Val Val Arg Leu Cys Arg Asp Leu Gly Tyr	
50 55 60	
Lys Gly Tyr Ser Asp Phe Arg Met Ala Leu Ala Val Asp Leu Ser Gln	
65 70 75 80	
Ser Ala Asn Gln Ser Gln Pro Lys Met Asp Gly Asp Ile Cys Glu Val	
85 90 95	
Ser Ala Gln Ser Ala Val Asp Ser Leu Met Asp Thr Ala Lys Leu Ile	
100 105 110	
Asp Arg Ala Ala Leu Asn Arg Ile Cys Glu Leu Val His Gly Ala Lys	
115 120 125	
Phe Ile Gly Cys Val Gly Val Gly Ala Ser Ser Ile Val Gly Arg Tyr	
130 135 140	
Leu Ala Tyr Arg Leu Val Arg Ile Gly Lys Lys Ala Ile Met Tyr Glu	
145 150 155 160	
Asp Thr His Leu Ala Ala Met Ser Ala Gly Gln Ser Val Val Gly Asp	
165 170 175	
Ala Trp Phe Ala Ile Ser Ser Gly Ser Thr Lys Glu Val Val His	
180 185 190	
Ala Ala Thr Gln Ala His Gln Arg Gly Val Pro Val Val Ser Leu Thr	
195 200 205	
Asn Ile Ser His Ser Pro Leu Ser Ser Ile Ser Asp Glu Met Leu Val	
210 215 220	
Ala Ala Arg Pro Glu Gly Pro Leu Thr Gly Gly Ala Phe Ser Ser Lys	
225 230 235 240	
Val Gly Ala Leu Leu Leu Val Asp Val Leu Ile Asn Thr Leu Leu Asp	
245 250 255	

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Val Tyr Pro Glu Tyr Ser Ala Ser Val Phe Gly Thr Ala Glu Val Ile  
 260                    265                    270

Leu Pro Leu Met Asp Ser  
 275

<210> SEQ\_ID NO 17  
 <211> LENGTH: 288  
 <212> TYPE: PRT  
 <213> ORGANISM: Haemophilus influenzae  
 <400> SEQUENCE: 17

Met Ala Lys Ser Gly Asn Val Leu Asn Lys Ile Gly Ser Leu Tyr Gln  
 1                    5                    10                    15

Ser Leu Thr Lys Ser Glu Lys Lys Ile Ala Asp Thr Ile Leu Arg Ser  
 20                    25                    30

Pro Asp Leu Val Ser Gln Cys Ser Leu Ser Glu Ile Ala Lys His Leu  
 35                    40                    45

Gln Val Gly Glu Ala Thr Leu Val Arg Phe Cys Arg Thr Ile Gly Phe  
 50                    55                    60

Lys Gly Phe Ser Glu Phe Lys Leu Glu Leu Ser Ile Glu Leu Ala Thr  
 65                    70                    75                    80

Lys Asp Asn Gln Asp Glu Glu Ile Leu Glu Thr Glx Ile Met Pro Glu  
 85                    90                    95

Asp Asp Glu Leu Thr Ile Ala Gln Lys Leu Gln Thr Ala Val Ala Asn  
 100                    105                    110

Val Met Glu Glu Thr Ile Asn Leu Leu Asp Leu Lys Gln Leu Glu Glu  
 115                    120                    125

Val Val Lys Val Leu Lys Lys Ala Arg Arg Ile Pro Leu Phe Cys Val  
 130                    135                    140

Cys Glu Glu Cys Val Thr Ala Glu Asp Ala Lys Asn Lys Leu Met Arg  
 145                    150                    155                    160

Ile Gly Pro Gln Val Val Ala Ser Gly Asn Asn His Pro Met Ala Met  
 165                    170                    175

Gln Ala Ala Leu Leu Thr Ser Ser Asp Val Ala Thr Gly Leu Ser Arg  
 180                    185                    190

Ser Gly Pro Ser Ala Glu Ile Ala His Ile Ile Lys Ile Ala Met Val  
 195                    200                    205

Asn Gly Ala Thr Thr Val Ala Leu Tyr His Ser Leu Met Ser Pro Val  
 210                    215                    220

Thr Glu Thr Ala Val Tyr Val Leu Val Asn Gly Asn Lys Val Gly Lys  
 225                    230                    235                    240

Leu Gln Gly Asp Ser Ile Gly Thr Lys Ile Ala Gln Leu Phe Val Leu  
 245                    250                    255

Asp Leu Ile Tyr Ala Leu Leu Val Gln Gly Glu Glu Asp Ile Ala Ala  
 260                    265                    270

Gln Thr Lys Gln Lys Thr Leu Asn Val Ile Leu Glu Gln Arg Ile Lys  
 275                    280                    285

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What is claimed is:

**1.** A method for screening a substance regulating the interaction between NanR and the transcriptional control region of a nan operon, comprising

(a) designing and displaying a tertiary structure of a complex of NanR protein and ManNAc-6P using the atomic coordinates of the complex shown in Table 3, wherein the complex comprises binding site amino acid residues R71, A137, S138, H163, S182, S183, S184, T187, E229, P231, G234 and K240;

(b) preparing candidate substances binding to NanR by employing the identified binding site amino acids from (a) to design and fit said substances; and

(c) examining binding affinity of the candidate substances for NanR and its regulation of the interaction between NanR and the transcriptional control region of nan operon,

wherein step (a) is carried out using a computer, and wherein the atomic coordinates of the complex shown in Table 3 are stored on non-transitory electronic media that is used by the computer.

**2.** The method according to claim 1, further comprising determining the candidate as a bacterial growth inhibitor, when the candidate binding to NanR has the NanR binding affinity similar to or higher than that of ManNAc-6P and maintains or increases interaction between NanR and the transcriptional control region of nan operon, compared to a control group treated without the corresponding candidate.

**3.** The method according to claim 1, further comprising determining the candidate as a bacterial growth stimulant, when the candidate binding to NanR has the NanR binding affinity similar to or higher than that of ManNAc-6P and decreases interaction between NanR and the transcriptional

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control region of nan operon, compared to a control group treated without the corresponding candidate.

**4.** A method for screening a substance regulating the interaction between NanR and ManNAc-6P, comprising

(a) designing and displaying a tertiary structure of a complex of NanR protein and ManNAc-6P using the atomic coordinates of the complex shown in Table 3, wherein the complex comprises binding site amino acid residues R71, A137, S138, H163, S182, S183, S184, T187, E229, P231, G234 and K240;

(b) preparing candidate substances binding to NanR by employing the identified binding site amino acids from (a) to design and fit said substances; and

(c) examining whether the candidate regulates the interaction between NanR protein and ManNAc-6P, wherein step (a) is carried out using a computer, and wherein the atomic coordinates of the complex shown in Table 3 are stored on non-transitory electronic media that is used by the computer.

**5.** The method according to claim 4, further comprising determining the candidate as a nan operon expression enhancer or as a bacterial growth stimulant, when the candidate increases interaction between NanR and ManNAc-6P, compared to a control group treated without the corresponding candidate.

**6.** The method according to claim 4, further comprising determining the candidate as a nan operon expression suppressor or as a bacterial growth inhibitor, when the candidate decreases interaction between NanR and ManNAc-6P, compared to a control group treated without the corresponding candidate.

\* \* \* \* \*